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(54) Title: TERMINAL COMPLEMENT INHIBITOR FU	ISION	GENES AND PROTEINS

(57) Abstract

Nucleic acid sequences encoding chimeric proteins that comprise a functional portion of a parent terminal complement inhibitor, such as CD59, and a heterologous transmembrane domain are provided. The parent terminal complement inhibitor is modified to inactivate its GPI signal sequence. The heterologous transmembrane domain serves to anchor the chimeric protein to the cell membrane without substantially interfering with the complement inhibitor activity of the terminal complement inhibitor. The nucleic acid sequences and encoded chimeric proteins can be used to protect cells from complement attack.

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TERMINAL COMPLEMENT INHIBITOR FUSION GENES AND PROTEINS

FIELD OF THE INVENTION

The present invention relates to terminal complement inhibitory proteins that have been genetically engineered to alter their attachment to the cell surface and to medical uses of such novel molecules.

BACKGROUND OF THE INVENTION

I. The Complement System

The complement system acts in conjunction with other immunological systems of the body to defend against intrusion of cellular and viral pathogens. There are at least 25 complement proteins, which are found as a complex collection of plasma proteins and membrane cofactors. The plasma proteins make up about 10% of the globulins in vertebrate serum. Complement components achieve their immune defensive functions by interacting in a series of intricate but precise enzymatic cleavage and membrane binding events. The resulting complement cascade leads to the production of products with opsonic, immunoregulatory, and lytic functions.

The lytic aspect of complement function is effected by the permeablization of target cell membranes as a direct action of an assemblage of complement proteins known individually as "terminal complement components" and, in their functional assemblage, as the membrane attack complex, or "MAC". (See Esser, 1991; and Bhakdi, et al., 1991.) The actions of the MAC, hereinafter referred to as "complement attack," create pores or leaky patches that lead to the disruption of osmotic and ionic gradients in target cells, which, at high enough MAC concentrations, causes cell death. Lower concentrations of MACs can produce other effects, including activation

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of endothelial cells and platelets. Inappropriate MAC activity can result in pathologic damage to cells and tissues.

The complement cascade progresses via the classical pathway or the alternative pathway. These pathways share many components, and, while they differ in their early steps, both converge and share the same terminal complement components responsible for complement attack and the activation and/or destruction of target cells.

The classical complement pathway is typically initiated by antibody recognition of and binding to an antigenic site on a target cell. The alternative pathway is usually antibody independent, and can be initiated by certain molecules on pathogen surfaces. Both pathways converge at the point where complement component C3 is cleaved by an active protease (which is different in each pathway) to yield C3a and C3b. Other pathways activating complement attack can act later in the sequence of events leading to various aspects of complement function, including the formation of the MAC.

induce anaphylatoxin that can is an degranulation of mast cells, resulting in the release of histamine and other mediators of inflammation. As opsonin, it binds to bacteria, multiple functions. viruses and other cells and particles and tags them for C3b can also form a removal from the circulation. complex with other components unique to each pathway to form classical or alternative C5 convertase, cleaves C5 into C5a (another anaphylatoxin), and C5b, which is the first of the terminal complement components that make up the MAC. (Amongst the several means by which complement attack can be initiated, proteolytic enzymes with relatively broad target protein specificities, including plasmin, elastase, and cathepsin G, can cleave C5 so as to mimic the action of C5 convertase and produce active C5b.) C5b combines sequentially with C6, C7, and C8 to form the C5b-8 complex at the surface of the target

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cell. Upon binding of several C9 molecules, the active MAC (C5b-9) is formed.

II. Regulation of the Complement System

Normally, the complement system is in a continuous state of spontaneous turnover. C3 can spontaneously acquire C3b functions, forming a functional C3 convertase and leading to the formation of more C3b. The C3b generated in this spontaneous fashion can also form C5 convertase and thus initiate the final steps in the cascade that forms the MAC.

Under normal conditions, blood flow will dilute and disperse the low levels of spontaneously activated complement components thus helping to prevent MAC buildup in any one location in the vasculature. In addition, homeostatic regulation of the actions of autologous complement proteins to prevent autoimmune attack is mediated by specific endogenous complement inhibitor proteins (CIPs), that can be found on the surfaces of most human cells. Ordinarily, blood flow and the action of CIPs suffice to render cells resistant to normal levels of spontaneous complement activation without injury or lysis. Under conditions of acute inflammation, and in various disease states where complement activation and MAC formation are accelerated, the normal quantity and activity of endogenous complement inhibitors may be inadequate to protect autologous cells from MAC-induced lysis and/or sublytic MAC-induced cell activation. Endogenous CIP activity may also be insufficient where there is stasis of the blood, and/or where there are defects in or deficiencies of naturally occurring inhibitors.

A number of CIPs have been identified that serve to protect cells from damage mediated by complement from concordant species. See Zalman, et al., 1986; Schonermark, et al., 1986; Nose, et al., 1990; and Sugita, et al., 1988. These inhibitors act at various defined points in the complement cascade. For example,

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CD55, also known as decay accelerating factor (DAF), exerts its major inhibitory effects on the actions of C3 convertase.

In cases where the complement cascade is initiated at points in the pathway after the C3 convertase step, such as through the generation of active C5b by broad spectrum proteases, DAF and other complement inhibitors acting at earlier steps in the cascade sequence are ineffective. There are, however, inhibitors that do not share this deficiency. These inhibitors act at the final steps in MAC assembly and thus can effectively block complement attack initiated by almost any means. These inhibitors are known as "terminal complement inhibitors" or "terminal CIPs."

15 III. <u>Terminal CIPs</u>

The most thoroughly characterized terminal CIP is the human protein CD59 (also known as "protectin", CD59 is a glycoprotein with an "MACIF", or "p18"). mass of 18-21 kilodaltons that apparent molecular protects cells from complement-mediated lysis. the outside of the cell by tethered to glycolipid moiety glycosyl-phosphatidylinositol (GPI) that anchors it in the cell membrane. CD59 is found associated with the membranes forming the surfaces of most human cells including erythrocytes, lymphocytes, and vascular endothelial cells. (See, for example, Sims, et al., U.S. Patent No, 5,135,916.)

CD59 appears to function by competing with C9 for binding to C8 in the C5b-8 complex, thereby decreasing the formation of the C5b-9 MAC (Rollins, et al., 1990). CD59 thus acts to reduce both cell stimulation and cell lysis by MACs (Rollins, et al., 1990; Rollins, et al., 1991; Stefanova, et al., 1989; Sugita, et al., 1988; Davies, et al., 1989; Holguin, et al., 1989; Okada, et al., 1989a; Meri, et al., 1990; Whitlow, et al., 1990; and Harada, et al., 1990). This activity of CD59 is for the most part species-selective, most efficiently

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blocking the formation of MACs under conditions where C8 and C9 are derived from homologous (i.e., human) serum (Venneker, et al., 1992). assimilation The purified CD59 into the plasma membrane of non-human erythrocytes (which are believed to be protected from homologous non-human complement attack by the action of their own cell surface complement inhibitor proteins) and oligodendrocytes (brain cells which are believed to be protected less, if at all, by cell surface proteins, but may be protected in vivo by the blood brain barrier) has shown that CD59 can protect these cells from cell lysis mediated by human complement. (Rollins, et al., 1990; Rollins, et al., 1991; Stefanova, et al., 1989; Meri, et al., 1990; Whitlow, et al., 1990; Okada, et al., 1989b; and Wing, et al., 1992).

cDNAs coding for CD59 have been cloned and the structure of the CD59 gene has been characterized (Davies, et al., 1989; Okada, et al., 1989b; Philbrick, et al., 1990; Sawada, et al., 1989; and Tone, et al., 1992). Non-human mammalian cells transfected with the cloned CD59 cDNA, and thereby expressing the human CD59 protein on their cell surfaces, have been shown to gain resistance to complement-mediated cell lysis (Zhao, et al., 1991; and Walsh, et al., 1991).

CD59 has been reported to be structurally related to the murine Ly-6 antigens (Philbrick, et al., 1990; and Petranka, et al., 1992). The genes encoding these antigens, also known as T-cell activating proteins, are members of the Ly-6 multigene family, and include Ly-6A.2, Ly-6B.2, Ly-6C.1, Ly6C.2, and Ly-6E.1. The gene encoding the murine thymocyte B cell antigen ThB is also a member of this family (Shevach, et al. 1989; and Gumley, et al., 1992).

A number of viral and non-human primate complement inhibitor proteins that are similar in structure and function to CD59 have been described (see Rother, et al., 1994; Albrecht, et al., 1992; commonly assigned,

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copending, U.S. patent application Serial No. 08/105,735, filed August 11, 1993, by William L. Fodor, Scott Rollins, Russell Rother, and Stephen P. Squinto, and entitled "Complement Inhibitor Proteins of Non-Human Primates;" and commonly assigned and copending PCT patent application Serial No. PCT/US93/00672, filed January 12, 1993, by Bernhard Fleckenstein and Jens-Christian Albrecht, and entitled "Complement Regulatory Proteins of Herpesvirus Saimiri".

These proteins -- BABCIP (SEQ ID NO:1), AGMCIP (SEQ ID NO:2), SQMCIP (SEQ ID NO:3), OWMCIP (SEQ ID NO:4), MARCIP (SEQ ID NO:5), and HVS-15 (SEQ ID NO:6) -- all share striking sequence homologies, including a distinctive conserved arrangement of cysteines within their amino acid sequences. These conserved patterns are most readily perceived by aligning the sequences of the proteins so that the cysteine residues are in register as seen in FIG. 1.

Cysteine residues of many proteins form a structural element referred to in the art as a "cysteine backbone". In proteins in which they occur, cysteine backbones play essential roles in determining the three-dimensional folding, tertiary structure, and ultimate function of the molecule. The proteins of the Ly-6 multigene family, as well as several other proteins, share a particular cysteine backbone structure referred to herein as the "Ly-6 motif". For example, the human urokinase plasminogen activator receptor (uPAR; Roldan, et al., 1990) and one of several squid glycoproteins of unknown function (Sgp2; Williams, et al., 1988) contain the Ly-6 motif.

Subsets of proteins having the Ly-6 motif can be identified by the presence of conserved amino acid residues immediately adjacent to the cysteine residues. Such conservation of specific amino acids within a subset of proteins can be associated with specific aspects of the folding, tertiary structure, and ultimate function of

the proteins. These conserved patterns are most readily perceived by aligning the sequences of the proteins so that the cysteine residues are in register.

discussed fully in the above-referenced, copending U.S. patent application Serial No. 08/105,735, the relevant portions of which are incorporated herein by reference, a series of non-human primate C5b-9 inhibitory proteins have been identified which are characterized by a cysteine backbone structure which defines a specific subset of the general Ly-6 motif.

Specifically, these non-human primate CIPs include polypeptides comprising a cysteine backbone with a Ly-6 motif characterized by the formula:

$$Cys-X_2-Cys-X_{6-9}-Cys-X_5-Cys-X_6-Cys-X_{12}-$$

 $Cys-X_5-Cys-X_{17}-Cys-X_0-Cys-X_4-Cys.$ (1)

In addition, the non-human primate C5b-9 inhibitory proteins include amino acid sequences conforming to the following formula:

Cys-X₅-(Thr or Ser)-Cys-X₁₁-(Gln or Arg)-

$$Cys-X_4-(Asn or Asp)-Cys-X_{17}-Cys-X_0-Cys-X_4-Cys.$$
 (2)

In both formulas, the X in X_n indicates a peptide containing any combination of amino acids, the n in Xn represents the length in amino acid residues of the peptide, and each X at any position can be the same as or different from any other X of the same length in any other position.

discussed fully in the above-referenced, copending, commonly assigned, PCT application Serial No. PCT/US 93/00672, the relevant portions of which are incorporated herein by reference, and in Albrecht, et al., 1992, a protein of the herpesvirus saimiri having C5b-9 inhibitory activity has been discovered (referred to herein as "HVS-15"). This viral protein has the Ly-6 motif which is characteristic of the non-human primate C5b-9 inhibitory proteins discussed above, i.e., its structure is described by formulas (1) and (2) above.

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In the discussion which follows, terminal CIPs comprising Ly-6 motifs are referred to as "Ly-6 terminal CIPs." These CIPs will in general satisfy formula (1) above and preferably also formula (2). Some variations, however, in the spacing between any two of the ten cysteines making up the Ly-6 motif and in the adjacent amino acids are to be expected in as yet uncharacterized terminal CIPs of other species.

Also, Petranka et al., 1993, and Norris, et al., 1993, have reported that in CD59 (SEQ ID NO:7) the disulfide bond between Cys6 and Cys13, as well as the disulfide bond between Cys64 and Cys69, can be disrupted by replacement of these cysteines with serines without substantially compromising the functionality of CD59. These cysteines correspond to the second, third, ninth, and tenth cysteines in the above formulas. Accordingly, as used herein, the term "Ly-6 terminal CIP" is intended to also include terminal complement inhibitor proteins conforming with the above formulas but with all or some of the second, third, ninth or tenth cysteines replaced with serine, or another amino acid.

IV. Other Cell Surface Complement Inhibitor Proteins

In addition to the Ly-6 terminal CIPs discussed above, other membrane bound CIPs have been described in the literature, including the following:

- (a) CD46 (membrane cofactor protein, MCP, see, for example, PCT patent publication No. WO 91/02002) is a 350 amino acid transmembrane (TM) protein found on all cells except red blood cells. CD46 binds to C3b, and, once bound, promotes the activity of proteases that cleave C3b into inactive fragments, thus preventing C3b accumulation on the cell surface and, in turn, protecting cells from complement attack. Both membrane bound and secreted forms of CD46 have been reported in the literature (Purcell et al., 1991).
- (b) CD55 (decay accelerating factor, DAF), mentioned above, is a GPI-anchored cell surface protein

present on all cells including red blood cells. Unlike CD46, CD55 does not destroy C3b. Rather, CD55 prevents C3b from reacting with other complement components, thus contravening complement mediated cytolysis. Both membrane bound and secreted forms of CD55 have been reported in the literature (Moran et al., 1992).

- (c) CD35 (complement receptor 1, CR1) is found on a select group of lymphocytes as well as erythrocytes, neutrophils, and eosinophils, and causes degradation of C3b molecules adhering to neighboring cells.
- (d) Factor H and C4b-binding protein, both of which inhibit alternative C3 convertase activity.

V. <u>Transplantation</u>

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Intrinsic activation of complement attack via the alternative pathway during storage of donor organs is responsible for certain problems associated with organ transplantation which arise as a result of endothelial cell stimulation and/or lysis by the C5b-9 MAC (Brasile, et al. 1985). Ex vivo complement attack leads to reduced vascular viability and reduced vascular integrity when stored organs are transplanted, increasing the likelihood of transplant rejection.

Ten percent of allogeneic transplanted kidneys with HLA-identical matches are rejected by in vivo immunologic (Brasile, et al. 1987). In 78% of the patients who reject organs under these conditions, cytotoxic antibodies binding to molecules on the surfaces of vascular endothelial cells are seen (Brasile, et al., Such antibody cytotoxicity is mediated by complement attack, and is responsible for the rejection of transplanted solid organs including kidneys and hearts (Brasile, et al., 1987; Brasile et al., 1985). Antibody primed, complement-mediated rejection is usually rapid and irreversible, a phenomenon referred to as hyperacute rejection.

In the xenogeneic setting, as when non-human organs are transplanted into human patients, activation of

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directed by antibodies attack complement molecules on the surfaces of endothelial cells lining the vessels of the donor organ is almost always observed. The prevalence of such xenoreactive antibodies accounts occurrence of hyperacute the nearly universal rejection of xenografts (Dalmasso, et al., 1992). world primates, including humans, have high levels of "natural" antibodies circulating preexisting determinants carbohydrate recognize predominantly expressed on the surface of xenogeneic cells discordant species. Recent evidence indicates that most of these antibodies react with galactose in an $\alpha 1-3$ linkage with galactose (Gal(α 1-3)Gal) (Sandrin, et al., 1993).

Old world primates lack the appropriate functional α -1,3-galactose transferase and thus do not express this carbohydrate epitope. Therefore, following transplantation of a vascularized xenogeneic donor organ, these high-titer antibodies bind to the Gal (α 1-3)Gal epitope on the vascular endothelium and activate the recipient's complement through the classical pathway. The massive inflammatory response that ensues from activation of the complement cascade leads to the destruction of the donor organ within minutes to hours.

exclusively antibodies are not Xenoreactive responsible for hyperacute rejection of discordant organs For example, erythrocytes from some in all cases. species can activate human complement via the alternative pathway and newborn piglets raised to be free almost xenografts antibodies reject preformed immediately. It is therefore likely that in some species combinations, activation of the alternative complement pathway contributes to graft rejection.

Endogenously-expressed, membrane-associated complement inhibitory proteins normally protect endothelial cells from autologous complement. However, the species restriction of complement inhibitors makes them

relatively ineffective with respect to regulating discordant xenogeneic serum complement. The lack of effective therapies aimed at eliminating this antibody and complement-mediated hyperacute rejection presents a major barrier to the successful transplantation of discordant animal organs into human recipients.

Recently, a report on a baboon-to-human liver transplant has been published in which the xenogeneic donor organ failed to exhibit signs of hyperacute rejection (Starzl, et al., 1993). The low levels of anti-baboon antibodies likely to be present in human blood make hyperacute responses less likely. However, it is believed that recently discovered baboon CIPs, which have been shown to be related to CD59 and to be effective against human complement, also played a role in maintaining the integrity of this xenotransplanted organ. (See U.S. patent application Serial No. 08/105,735, referred to above.)

The lack of hyperacute rejection seen in the baboon to human xenotransplant discussed above suggests that complement inhibitor proteins effective against human complement may, in combination with other anti-rejection strategies, allow safe and effective xenotransplantation of transgenic animal organs expressing such proteins into human patients.

VI. GPI-Anchored CIPs and Modifications Thereof

GPI-anchored terminal CIPs share certain properties that make them less desirable than transmembrane (TM) proteins for use as complement inhibiting agents for the protection of transplanted cells or organs.

GPI-anchored terminal CIPs, including CD59, BABCIP, and AGMCIP, can be cleaved from cell surfaces by specific phospholipase enzymes that hydrolyze GPI anchors. Such phospholipases are present in the serum (phospholipase D, Davitz, et al., 1987), and may also be released from cells in response to ischemia (phospholipase C, Vakeva, et al., 1992). Since ischemia is an unavoidable

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concomitant of transplantation, the process of transplantation may serve to remove native and/or artificially introduced GPI-anchored terminal CIPs from the very cells within the transplanted organ that they are meant to protect.

Another mechanism by which GPI-anchored proteins are removed from the cell surface is the incorporation of such proteins into membrane vesicles and the subsequent the vesicles from the cell. of shedding vesiculation can occur in response to various stimuli, such as ischemia-induced complement attack. It has been reported that GPI-anchored proteins are concentrated in these vesicles relative to their concentration in the cell membrane, a phenomenon that may reflect involvement of these proteins in the vesiculation process itself (Butikofer, et al., 1989; Brown, et al., 1992; Whitlow, et al., 1993). Such preferential incorporation into shed vesicles can reduce the concentrations of GPI-anchored cell surface, including the proteins onthe GPI-anchored terminal CIPs. Such concentrations reductions of terminal CIP concentrations, particularly in response to complement attack, may occur at just those times when inhibition of complement is most needed.

In addition to their susceptibility to removal from the cell surface, GPI-anchored proteins also suffer from the problem that their production may be limited in various cell types. That is, only so many GPI-anchored molecules can normally be produced by a cell within a given time frame, so that introducing genes for further GPI-anchored proteins may not in fact result in substantial increases in the amount of protein actually present on the cell surface.

The limiting case of this problem involves cells which are incapable of producing any GPI-anchored proteins. The clinical disease of paroxysmal nocturnal hemoglobinuria (PNH) involves cells of this type, specifically, blood cells which do not produce GPI-

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anchored terminal CIPs. As discussed in copending, commonly assigned, U.S. patent application Serial No. 08/206,189, entitled "Method for the Treatment of Paroxysmal Nocturnal Hemoglobinuria", which is being filed concurrently herewith in the names of Russell Rother, Scott A. Rollins, Seth A. Fidel, and Stephen P. Squinto, PNH cells can be made resistant to complement attack through the use of the transmembrane terminal CIPs described herein.

A further drawback of GPI-anchored proteins involves the ability of these proteins to transduce signals into the cell upon being cross-linked by specific antibodies and presumably upon binding their natural ligand (Okada, et al., 1989b; Seaman, et al., 1991; Su, et al., 1991; Deckert, et al., 1992; Cinek, et al., 1992; Card, et al., 1991; Groux, et al., 1989; and Stefanova, et al., 1991). Possible undesirable cellular responses intracellular signals can include phospholipase activation and/or release, and the stimulation of vesicle formation and shedding, both of which, above, can result in the loss of GPI-anchored proteins from the cell surface. Thus, the very GPI-anchored terminal CIPs that are used to protect the cells of a transplanted organ from complement attack may activate the cellular events that lead to their removal from the cell surface.

Work has been performed in which the means of attachment of GPI-anchored proteins to the outer cell surface has been varied from their natural GPI anchors by substitution of other anchoring moieties (Su, et al., 1991; and Lublin, et al., 1991).

For example, chimeric derivatives of CD55, containing amino acids 1-304 of CD55 fused to a fragment of CD46 which includes the protein's transmembrane domain (i.e., amino acids 270-350 of CD46) or to a fragment of the human major histocompatibility protein HLA-B44 which includes its transmembrane domain (i.e., amino acids

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262-338 of HLA-B44), have been reported to retain levels of function equivalent to native CD55 (Lublin, et al., 1991). Significantly, with regard to the present invention, no such substitutions have been made with terminal CIPs and no such molecules have been developed for clinical use and, in particular, for use in constructing transgenic organs for transplantation.

VII. Protein Structure and Function

Minor alterations of protein primary structures (amino acid sequences) can have profound effects on their functional properties. The best known example of this phenomenon is in the case of sickle cell anemia, in which a single amino acid alteration, namely, a change in residue 6 of the beta chain of hemoglobin from Glu to Val, is sufficient to change the oxygen binding properties of the hemoglobin molecule and to thereby cause sickle cell disease.

The insertion of heterologous amino acid sequences representing new domain structures into a protein can also have significant effects on the protein's functional properties. For example, the introduction of a 10 amino acid epitope of the c-myc proto-oncogene (known as the myc tag) to the <u>int-1</u> proto-oncogene alters functional properties of int-1. Specifically, C57MG mammary epithelial cells are transformed by wild-type int-1, but not by the myc-tagged int-1, while residual function of the myc-tagged int-1 gene is seen in a more examining effects sensitive assay on Drosophila development (McMahon et al., 1989).

Additionally, substitution of homologous sequences from heterologous proteins can have profound effects on protein function. For example, replacement of either of the two most carboxyl-terminal 12 amino acid segments of the mouse nerve growth factor gene with homologous segments from the related mouse brain derived neurotrophic factor gene reduces the activity of the molecule by 50%. That is, the carboxyl-terminal region is

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particularly sensitive to substitution with a homologous sequence from a heterologous protein, such a substitution having sufficient impact on protein function to decrease activity by 50%. A similar decrease in activity is seen following substitution of the amino terminus (Suter, et al., 1992).

All Ly-6 terminal CIPs are believed to share the property of being attached to cell membranes by means of a GPI linkage. As understood in the art, the addition of such a GPI moiety to a nascent protein coincides with a proteolytic processing step that removes a number of amino acid residues from the carboxyl-terminus of the polypeptide. Accordingly, mature Ly-6 terminal CIPs do not include all of the amino acids specified by the full nucleic acid molecules that encode Specifically, they do not include some or all of the amino acid residues downstream of the cysteine backbone Ly-6 motif, e.g., the amino acids downstream of cysteine 69 of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7 (CD59) and downstream of cysteine 72 of SEO ID NO:3. (As used herein. "downstream" means towards the carboxyl terminus of the polypeptide or towards the 3' end of the coding strand of the nucleic acid molecule coding for the polypeptide and "upstream" means towards the amino terminus of the polypeptide or towards the 5' end of the coding strand of the nucleic acid molecule coding for the polypeptide.) It is not known which amino acids downstream of the Ly-6 cysteine backbone motif are present or absent in any of these terminal CIPs when they are in the mature, GPI anchored state.

As discussed in detail below, the present invention involves the removal of selected amino acids of such Ly-6 terminal CIPs downstream of the Ly-6 motif. In view of the foregoing state of the art, it was not known, prior to the present invention, what effects such amino acid removal would have on terminal CIP function. In

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particular, it was not known if Ly-6 terminal CIPs would retain any complement inhibitory activity after such removal.

Various attempts have been made to examine the effects of GPI anchors on protein function. In the case of CD55, the substitution of protein fragments that contain a transmembrane domain for the carboxyl-terminal sequences believed to be involved in the addition of the GPI anchor (referred to hereinafter as the "GPI signal sequence") results in a protein with equal activity to the native GPI-anchored protein (Lublin, et al., 1991). In the case of the Ly-6 protein, Ly-6E (Ly-6E.1), which surface protein that GPI-anchored cell structurally related to CD59 (Philbrick, et al. 1990), the substitution of a fragment containing a transmembrane domain for the carboxyl-terminal GPI signal sequences downstream of the Ly-6 motif produces a non-functional protein, i.e., a protein which is not capable of activating T-cells (Su, et al., 1991).

SUMMARY OF THE INVENTION

In view of the foregoing, it is an object of this invention to provide novel proteins that can be used in controlling the complement system of humans and other animals. It is a further object of the invention to provide nucleic acid sequences and associated genetic engineering constructs for producing such proteins either in vitro or in vivo.

More particularly, it is the object of the invention to provide novel proteins that are Ly-6 terminal complement inhibitors, but are anchored to the cell surface by means independent of GPI anchoring. It is an additional object of the invention to provide molecules of this type that will not transmit an activating signal into the cells to which they are bound, e.g., endothelial cells, lymphocytes, or platelets, either after antibody crosslinking, or upon binding of the terminal CIP to its ligand. It is a further object of the invention to

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provide molecules of this type that cannot be removed from the surfaces of the cells to which they are bound by the actions of lipid cleaving enzymes such as phospholipases and which are not preferentially incorporated into shed vesicles.

To achieve the foregoing and other objects, the present invention, in accordance with certain of its aspects, provides the complete cDNA sequences of chimeric genes encoding chimeric protein products which comprise the fusion of a Ly-6 terminal CIP with a heterologous transmembrane (TM) domain. Prior to fusion, selected amino acid residues located downstream from the Ly-6 motif of the terminal CIP are deleted. The invention also comprises the chimeric protein products encoded by these genes, such chimeric molecules being referred to hereinafter as TMTCIPs (i.e., transmembrane terminal complement inhibitor proteins). In the preferred embodiments of the invention, the chimeric proteins have greater than 50% of the complement inhibitory activity of the native, GPI-anchored terminal CIP from which the TMTCIP is derived where such activity is preferably measured using a dye release assay of the type described below in Example 4.

The protection from complement attack offered by the TMTCIPs of the invention can be provided via gene transfer for the therapeutic prevention of pathologic complement attack in, for example, transplantation. In a preferred form of such therapy, the expression of the TMTCIP can be directed to the surfaces of cells of non-human animal organs, e.g., organs of non-human transgenic animals, in order to protect such organs from complement attack upon transplantation into a human patient.

The accompanying drawings, which are incorporated in and constitute part of the specification, illustrate certain aspects of the preferred embodiments of the invention and, together with the description, serve to

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explain certain principles of the invention. It is to be understood, of course, that both the drawings and the description are explanatory only and are not restrictive of the invention.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 shows aligned amino acid sequences of Human, African Green Monkey, Baboon, Owl Monkey, Marmoset, Squirrel Monkey, and Herpesvirus Saimiri Ly-6 terminal CIPs (CD59, AGMCIP, BABCIP, OWMCIP, MARCIP, SQMCIP, and HVS-15, respectively). The cysteine residues making up the Ly-6 cysteine backbone motif of each protein are underlined.

FIG. 2 shows a comparison of the cell surface expression of CD59 epitopes on Balb/3T3 cells. The three traces represent cell surface expression profiles of a positive Balb/3T3 clone expressing the CD59-MCP TMTCIP (CD59-TM), a native human CD59 transfectant (CD59-GPI) as a positive control and a vector (pcDNA3, Invitrogen, San Diego, CA) without insert transfectant (Vector Control) as a negative control.

FIG. 3 shows a comparison of the cell surface expression of CD59 epitopes on mouse L cells. The broad trace represents cell surface expression profiles of pooled L cells transduced with retroviral virion particles generated using the pL-CD59-MCP-TM-SN vector (CD59-TM). Also shown are profiles of pooled L cells transduced either with retroviral virion particles generated using the pL-CD59-GPI-SN vector (CD59-GPI) or with retroviral virion particles generated using the pLXSN vector with no insert (Vector Control), as negative controls.

FIG. 4 shows cell surface levels of CD59 antigens on stably transfected Balb/3T3 cells before and after PI-PLC digestion. FIG. 4A shows data obtained using a clone expressing the native human CD59 molecule (CD59-GPI). FIG. 4B shows data obtained using a clone expressing the CD59-MCP TMTCIP (CD59-TM). In each panel, the traces

WO 95/23512 PCT/US95/02944

-19-

labeled "A" and "B" represent cells stained with the secondary antibody alone, without or with PI-PLC treatment, respectively. In each panel, the traces labeled "C" and "D" represent cells stained with both the primary (CD59 specific) antibody and the secondary antibody with or without PI-PLC treatment, respectively.

FIG. 5 shows data obtained from dye release assays performed using the transfected Balb/3T3 cells employed in obtaining the data of FIG. 2 and FIG. 4. The cells were challenged with 20% human C8 depleted serum supplemented with a mixture of equal parts of purified human C8 and C9. The amounts, in micrograms per milliliter final concentration, of the mixture of human C8 and C9 added are indicated on the abscissa and the percent of dye release is indicated on the ordinate.

FIG. 6 shows data obtained from dye release assays performed using the transfected mouse L cells employed in obtaining the data of FIG. 3. The cells were challenged with 20% human C8 depleted serum supplemented with a mixture of equal parts of purified human C8 and C9. The amounts, in micrograms per milliliter final concentration, of the mixture of human C8 and C9 added are indicated on the abscissa and the percent of dye release is indicated on the ordinate.

25 <u>DESCRIPTION OF THE PREFERRED EMBODIMENTS</u>

As discussed above, the present invention provides the complete cDNA sequences of chimeric genes encoding chimeric protein products which comprise the fusion of a Ly-6 terminal CIP and a heterologous transmembrane domain.

I. <u>Terminal CIPs</u>

A variety of terminal CIPs can be used in the practice of the invention. In particular, Ly-6 terminal CIPs can be used. In addition to sharing the homologies shown in formulas (1) and (2) above, the Ly-6 terminal CIPs also share a variety of other homologies which can be seen in the aligned amino acid sequences of FIG. 1.

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Homologies found downstream from the Ly-6 cysteine motif of these terminal CIPs include an N immediately following the last C of the Ly-6 motif, another N 6-8 residues downstream from the last C of the Ly-6 motif (referred to herein as the "truncation-Asn"), and the following consensus sequence, hereinafter referred to as the "downstream consensus sequence" which includes the aforementioned truncation-Asn at the third position in the sequence:

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(Lor I) (E or K) N (G or I) (G or K) (T or R) (S or T) (Lor I) S (K or E or D) K (T or A) (V or I or L) (Lor V) L L (V or L) (A or T or I) (P or L) (F or L) L (A or V) (A or T) A W (S or C or N) (L or R or F) (H or P) (P or L).

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In addition to these structural commonalities, testing of the Ly-6 terminal CIPs of FIG. 1 has shown that they share the ability to substantially inhibit the (See U.S. activity of human complement. 08/105,735, and PCT application Serial No. PCT/US93/00672, referred application Serial No. above.) In particular, each of CD59, AGMCIP, BABCIP, and HVS-15 had substantial SOMCIP, OWMCIP, complement inhibitory activity. MARCIP was not tested, but is also expected to have such activity.

25 II. <u>Transmembrane Domains</u>

As known in the art, transmembrane proteins may span the membrane once or several times along the length of There are in general two their amino acid chains. different ways in which a transmembrane protein that spans the membrane only once may be embedded in a Most commonly, these proteins have their membrane. located towards single transmembrane domain carboxyl-terminal end of the polypeptide chain and are oriented so that the region amino-terminal to the transmembrane domain is outside the cell or non-cytoplasmic cellular compartment and the region carboxyl-terminal to the transmembrane domain is in the cytoplasmic compartment. The second orientation of a transmembrane protein with a single membrane spanning transmembrane domain is the opposite of this common arrangement, that is, the region amino-terminal to the transmembrane domain is in the cytoplasmic compartment and the region carboxyl-terminal to the transmembrane domain is located outside the cell or in a non-cytoplasmic cellular compartment.

Other transmembrane proteins cross the membrane several times. Most commonly, eukaryotic representatives of this type of transmembrane protein have seven consecutive transmembrane domains, most of them connected by short hydrophilic loop regions.

Transmembrane proteins in general include at least one contiguous stretch of amino acid residues which resides in the lipid bilayer membrane (referred to hereinafter as "membrane amino acids"), and at least two contiguous stretches of amino acid residues which extend away from the membrane, one generally cytoplasmic (referred to hereinafter as "cytoplasmic amino acids"), and one generally extracellular or sequestered in a noncytoplasmic cellular compartment (referred to hereinafter as "extracellular amino acids"). As referred to herein, cytoplasmic amino acids and extracellular amino acids always include at least one charged amino acid residue immediately adjacent to the membrane amino (referred to herein as the "first cytoplasmic amino acid" and the "first extracellular amino acid," respectively).

Membrane amino acids are characterized as groups of at least about 20 amino acids (the minimum generally needed to span a membrane), most of which are hydrophobic (uncharged) amino acids. Charged (hydrophilic) amino acids are usually absent from these groups, but in some cases two hydrophilic residues of opposite charge may lie close together inside the membrane where they neutralize each other.

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Transmembrane domains derived from a variety of transmembrane proteins can be used in the practice of the transmembrane domains invention. However, cytoplasmic amino acids which include cysteine residues in close proximity to the first cytoplasmic amino acid may be expressed at lower levels on the cell surface than transmembrane domains that do not contain such cysteines. This decreased expression is believed to result from the cysteine residues propensity of these intermolecular bonds with similarly placed cysteines of adjacent nascent transmembrane protein molecules. intermolecular cysteine linkages cause aggregation of the nascent transmembrane proteins, generally within the Golgi apparatus (where newly synthesized transmembrane proteins are processed within the typical cell), and thus block the transport of such nascent proteins to the cell surface.

the TMTCIP molecules to the With regard invention, it is notable that transfection of mammalian cells with an expression vector encoding a chimeric terminal CIP containing a putative transmembrane domain from the herpesvirus saimiri CCPH gene (see PCT patent application Serial No. PCT/US93/00672, mentioned above) does not result in high enough levels of cell surface expression of terminal CIP epitopes to be detected by FACS analysis (see Example 1). The cytoplasmic amino acids of this putative transmembrane domain include cysteine residues spaced two and five amino acids from the first cytoplasmic amino acid, a histidine. presence of these cysteines is believed to be responsible for the low levels of expression seen with this putative transmembrane domain. For this reason, transmembrane domains of this type are not preferred for use with the present invention.

As used herein, the term "transmembrane domain" is intended to comprise: 1) the portion of a transmembrane protein which spans the membrane, i.e., the at least

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about twenty membrane amino acids normally required for this purpose, 2) the adjacent predominantly charged cytoplasmic amino acids within about five to about ten residues from the membrane amino acids, and 3) the adjacent predominantly charged extracellular amino acids within about five to about ten residues from the membrane amino acids. These adjacent predominantly charged cytoplasmic and extracellular amino acids are involved in anchoring the protein in the membrane. As discussed above, preferred transmembrane domains do not include cytoplasmic amino acids that are cysteine residues within five amino acids of the first cytoplasmic amino acid.

While it is possible to examine a protein sequence and pick out a region with about 20 consecutive hydrophobic amino acids, some transmembrane domains, as discussed above, contain a small number of hydrophilic amino acids interspersed within their predominantly hydrophobic residues. Accordingly, transmembrane domains are more effectively identified by using hydrophobicity scales to compute hydropathy plots (Branden, et al., 1991).

Hydrophobicity scales provide a numerical value for the hydrophobicity of individual amino acids. These scales have been developed on the basis of solubility measurements of amino acids in different solvents, vapor pressures of side-chain analogues, analysis of side-chain distributions within soluble proteins, and theoretical energy calculations (Kyte, et al., 1982; and Engelman, et al., 1986).

Hydropathy plots are computed from amino acid sequences using hydrophobicity values as follows. First, for each position in the sequence, a hydropathic index is calculated. The hydropathic index is the mean value of the hydrophobicity of the amino acids within a "window," usually 19 residues long, around each position. The hydropathic indices are then plotted versus amino acid sequence position to produce the hydropathy plot.

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Transmembrane domains are then identified from the hydropathy plots by searching for regions where the hydropathic index is high for a number of consecutive positions in the sequence, e.g., by searching for regions with broad peaks with high positive (i.e., hydrophobic) values.

In terms of the present invention, the transmembrane domain will preferably have a hydropathic index greater than about +0.5, using the scale of Kyte et al. (Kyte et al., 1982) and a window of 19 amino acids, over a region of at least about 12 amino acid residues.

Additional contiguous amino acids of the transmembrane protein can be included in or encoded by the chimeric molecules of the invention provided those additional amino acids do not substantially impair the insertion of the transmembrane domain into the membrane, the transport of the nascent chimeric protein to the cell surface, or the complement inhibitory activity of the terminal CIP portion of the chimeric molecule.

While the molecules of the present invention may be constructed with any functional transmembrane domain, one derived from a protein with only a single transmembrane domain and having the region carboxyl-terminal to its transmembrane domain in the cytoplasm is preferred. large number of such proteins have been reported in the literature, including the following: CD46; the major histocompatibility antigens and related transmembrane proteins of the immunoglobulin multigene superfamily including intercellular adhesion molecules, ICAM-1 (CD54), ICAM-2, ICAM-3, VCAM-1, PECAM-1 (CD31) and (CD44); the selectins, including E-selectin, HCAM L-selectin, and P-selectin (CD62); the Alzheimer's amyloid precursor protein; the insulin receptor; the epidermal growth factor receptor; the gp41 protein of the AIDs virus, HIV; the p21 proteins of HTLV1 and HTLV2; and the p15E proteins of the murine and feline leukemia viruses.

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TM domains derived from any of these proteins, as well as from other transmembrane proteins, can be used in the practice of the invention. These domains can be most easily used by incorporating into the chimeric molecule the entire carboxyl end of the transmembrane protein beginning at a point upstream from the transmembrane A particularly preferred TM domain is that constituting amino acids 294 to 326 of CD46 (MCP, SEQ ID This domain can be conveniently used along with amino acids 327 to 350, which comprise the carboxyl end of the CD46 protein downstream from the transmembrane domain of this molecule, and along with amino acids 270 to 293 upstream of the TM domain which do not interfere with insertion of the CD46 TM domain into cell membranes and, as shown below, do not inhibit the complement inhibitory activity of Ly-6 terminal CIPs.

In addition to using hydropathy plots to identify TM domains suitable for use in the present invention, such domains can also be identified biochemically using, for example, protease digestion techniques or by making chimeric molecules containing soluble proteins operatively linked to signal sequences and containing putative transmembrane domains, and assaying for membrane insertion of the chimeric protein.

25 III. TMTCIP Genes and Vectors Containing Such Genes

The isolation, truncation, and fusion of the nucleic acid fragments encoding the terminal CIP and the TM domain are performed using recombinant nucleic acid techniques known in the art, including: PCR generation of the desired fragments and/or restriction digestion of cloned genes; PCR fusion of the desired fragments; or enzymatic ligation of restriction digestion products (Sambrook, et al., 1989; and Ausubel et al., 1992). Alternatively, the nucleic acid molecules encoding the TMTCIPs of the invention or any or all of the nucleic acid fragments used to assemble the chimeric genes for

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the TMTCIPs can be synthesized by chemical means (Talib, et al., 1991).

The chimeric genes of the invention are prepared by 1) truncating the nucleic acid sequence for a Ly-6 terminal CIP so as to remove selected amino acid residues downstream of the Ly-6 motif in order to inactivate the normal GPI signal sequence, and 2) fusing the truncated sequence to a sequence coding for a selected TM domain and desired amino acids surrounding the TM domain.

The truncation of the nucleic acid sequence encoding the Ly-6 terminal CIP will remove at least some of the carboxyl-terminal amino acid residues downstream from the Asn which is located between 6 and 8 amino acid residues after the last (tenth) Cys of the Ly-6 motif. This Asn is also located at the third position in the downstream consensus sequence presented above, i.e., it is the truncation-Asn defined above. All known Ly-6 terminal CIPs include such a truncation-Asn.

In some cases, all of the amino acid residues after the truncation-Asn are removed. Alternatively, less than all can be removed, the criterion being that sufficient numbers of residues are removed so that the GPI signal sequence is inoperative. In general, the simplest approach is to remove all amino acid residues downstream of the truncation-Asn. If desired, the truncation can extend further upstream from the truncation-Asn. preferably starting at a point downstream from the last Cys of the Ly-6 motif. Truncations beginning upstream from the last Cys of the Ly-6 motif are in general not preferred, but can be used if desired. The criterion for truncations upstream of the truncation-Asn is the requirement that the TMTCIP has greater than 50% of the complement inhibitory activity of the parent (native) Ly-6 terminal CIP.

In terms of the Ly-6 terminal CIPs of FIG. 1, the preferred truncation comprises all of the amino acids downstream of Asn 77 of BABCIP (SEQ ID NO:1), Asn 75 of

AGMCIP (SEQ ID NO:2), Asn 80 of SQMCIP (SEQ ID NO:3), Asn 77 of OWMCIP (SEQ ID NO:4), Asn 77 of MARCIP (SEQ ID NO:5), Asn 77 of HVS-15 (SEQ ID NO:6), and Asn 77 of CD59 (SEQ ID NO:7). Of these Ly-6 terminal CIPs, CD59 is preferred. As discussed above, a preferred TM domain is from CD46. Accordingly, a particularly preferred embodiment of the invention comprises residues 1-77 of CD59 (SEQ ID NO:7) fused to amino acids 270-350 of CD46 (SEQ ID NO:8)

In addition to the foregoing, the present invention provides recombinant expression vectors which include nucleic acid fragments encoding the chimeric TMTCIPs of the invention. The nucleic acid molecule coding for such a chimeric protein can be inserted into an appropriate expression vector, i.e., a vector that contains the necessary elements for the transcription and translation of the inserted protein-encoding sequence. The necessary transcriptional and translational signals can also be supplied by the genes used to construct the fusion genes of the invention and/or their flanking regions.

transcriptional and translational sequences for expression vector systems to be used to direct expression in vertebrate cells may be provided by viral sources. For example, commonly used promoters and enhancers are derived from Polyoma virus, Adenovirus, Simian Virus 40 (SV40), the Molony murine leukemia virus (MMLV), including the long terminal repeat (MMLV-LTR), cytomegalovirus (CMV), including immediate-early gene 1 promoter and cytomegalovirus enhancer. Retroviral expression vectors are a preferred system for expression of the TMTCIPs of the invention.

The manipulation of retroviral nucleic acids to construct retroviral vectors and packaging cells is accomplished using techniques known in the art. See Ausubel, et al., 1992, Volume 1, Section III (units 9.10.1 - 9.14.3); Sambrook, et al., 1989; Miller, et al., 1989; Eglitis, et al., 1988; U.S. Patents Nos. 4,650,764,

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4,861,719, 4,980,289, 5,122,767, and 5,124,263; as well as PCT Patent Publications Nos. WO 85/05629, WO 89/07150, WO 90/02797, WO 90/02806, WO 90/13641, WO 92/05266, WO 92/07943, WO 92/14829, and WO 93/14188.

In particular, the retroviral vectors of the invention can be prepared and used as follows. First, a TMTCIP retroviral vector is constructed and packaged into non-infectious transducing viral particles (virions) using an amphotropic packaging system, preferably one suitable for use in gene therapy applications.

Examples of such packaging systems are found in, for example, Miller, et al., 1986; Markowitz, et al., 1988; Cosset, et al., 1990; U.S. Patents Nos. 4,650,764, 4,861,719, 4,980,289, 5,122,767, and 5,124,263, and PCT Patent Publications Nos. WO 85/05629, WO 89/07150, WO 90/02797, WO 90/02806, WO 90/13641, WO 92/05266, WO 92/07943, WO 92/14829, and WO 93/14188. A preferred packaging cell is the PA317 packaging cell line (ATCC CRL 9078).

The generation of "producer cells" is accomplished by introducing retroviral vectors into the packaging cells. Examples of such retroviral vectors are found in, for example, Korman, et al., 1987; Morgenstern, et al., 1990; U.S. Patents Nos. 4,405,712, 4,980,289, and 5,112,767; and PCT Patent Publications Nos. WO 85/05629, WO 90/02797, and WO 92/07943. A preferred retroviral vector is the MMLV derived expression vector pLXSN (Miller, et al., 1989). The retroviral vector used in the practice of the present invention will be modified to include the chimeric gene encoding the TMTCIP.

The producer cells generated by the foregoing procedures are used to produce the retroviral vector particles (virions). This is accomplished by culturing of the cells in a suitable growth medium. Preferably, the virions are harvested from the culture and administered to the target cells which are to be transduced, e.g., xenogeneic cells to be used for

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transplantation into a patient whose complement can be inhibited by the Ly-6 terminal CIP of the TMTCIP, cells of a xenogeneic organ to be used for transplantation into such a patient, the patient's own cells, and other cells to be protected from complement attack, as well as stem cells such as embryonic stem cells, which can be used to generate transgenic cells, tissues, or organs for transplantation. Alternatively, when practicable, the target cells can be co-cultured with the producer cells. Suitable buffers and conditions for stable storage and subsequent use of the virions can be found in, for example, Ausubel, et al., 1992.

Pharmaceutical compositions containing the retroviral vector particles of the invention can be administered in a variety of unit dosage forms. The dose will vary according to, e.g., the particular vector, the manner of administration, the particular disease being treated and its severity, the overall health condition and age of the patient, the condition of the cells being treated, and the judgment of the physician. Dosage levels for transduction of mammalian cells are generally between about 10^6 and 10^{14} colony forming units of retroviral vector particles per treatment.

A variety of pharmaceutical formulations can be used for administration of the retroviral vector particles of the invention. Suitable formulations are found in, for Remington's Pharmaceutical Sciences, example, Publishing Company, Philadelphia, PA, 17th ed., 1985, and will include a pharmaceutically effective carrier, such as saline, buffered (e.g., phosphate buffered) saline, Hank's solution, Ringer's solution, dextrose/saline, glucose solutions, and the like. The formulations may contain pharmaceutically acceptable auxiliary substances as required, such as, tonicity adjusting agents, wetting agents, bactericidal agents, preservatives, stabilizers, and the like.

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IV. Transgenic Animals

In accordance with certain aspects of the invention, the nucleic acid molecules of the present invention are used to generate engineered transgenic animals example, rodent, e.g., mouse, rat, capybara, and the like, lagomorph, e.g., rabbit, hare, and the like, ungulate, e.g., pig, cow, goat, sheep, and the like, etc.) that express the TMTCIPs of the invention on the surfaces of their cells (e.g., endothelial cells) using techniques known in the art. These techniques include, are not limited to, microinjection, e.q., pronuclei, electroporation of ova or zygotes, nuclear and/or the stable transfection or transplantation, transduction of embryonic stem cells derived from the animal of choice.

A common element of these techniques involves the preparation of a transgene transcription unit. Such a unit comprises a DNA molecule which generally includes:

1) a promoter, 2) the nucleic acid sequence of interest, i.e., the sequence encoding the TMTCIP of the present invention, and 3) a polyadenylation signal sequence. Other sequences, such as, enhancer and intron sequences, can be included if desired. The unit can be conveniently prepared by isolating a restriction fragment of a plasmid vector which expresses the TMTCIP protein in, for example, mammalian cells. Preferably, the restriction fragment is free of sequences which direct replication in bacterial host cells since such sequences are known to have deleterious effects on embryo viability.

The most well known method for making transgenic animals is that used to produce transgenic mice by superovulation of a donor female, surgical removal of the egg, injection of the transgene transcription unit into the pro-nuclei of the embryo, and introduction of the transgenic embryo into the reproductive tract of a pseudopregnant host mother, usually of the same species. See Wagner, U.S. Patent No. 4,873,191, Brinster, et al.,

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1985, Hogan, et al., 1986, Robertson 1987, Pedersen, et al., 1990.

The use of this method to make transgenic livestock is also widely practiced by those of skill in the art. As an example, transgenic swine are routinely produced by the microinjection of a transgene transcription unit into pig embryos. See, for example, PCT Publication No. W092/11757 In brief, this procedure may, for example, be performed as follows.

First, the transgene transcription unit is gel isolated and extensively purified through, for example, an ELUTIP column (Schleicher & Schuell, Keene, NH), dialyzed against pyrogen free injection buffer (10mM Tris, pH7.4 + 0.1mM EDTA in pyrogen free water) and used for embryo injection.

Embryos are recovered from the oviduct hormonally synchronized, ovulation induced preferably at the pronuclear stage. They are placed into a 1.5 ml microfuge tube containing approximately 0.5 ml of embryo transfer media (phosphate buffered saline with 10% fetal calf serum). These are centrifuged for 12 minutes at 16,000 x g in a microcentrifuge. Embryos are removed from the microfuge tube with a drawn and polished Pasteur pipette and placed into a 35 mm petri dish for examination. If the cytoplasm is still opaque with lipid such that the pronuclei are not clearly visible, the embryos are centrifuged again for an additional 15 Embryos to be microinjected are placed into a drop of media (approximately 100 μ l) in the center of the lid of a 100 mm petri dish. Silicone oil is used to cover this drop and to fill the lid to prevent the medium from evaporating. The petri dish lid containing the embryos is set onto an inverted microscope equipped with both a heated stage (37.5-38°C) and Hoffman modulation contrast optics (200X final magnification). A finely drawn and polished micropipette is used to stabilize the embryos while about 1-2 picoliters of injection buffer

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containing approximately 200-500 copies of the purified transgene transcription unit is delivered into the nucleus, preferably the male pronucleus, with another finely drawn and polished micropipette. surviving the microinjection process as judged by morphological observation are loaded into a polypropylene transfer into the recipient for ID) (2 mm pseudopregnant sow.

Offspring are tested for the presence of the transgene by isolating genomic DNA from tissue removed from the tail of each piglet and subjecting about 5 micrograms of this genomic DNA to nucleic acid hybridization analysis with a transgene specific probe.

Another commonly used technique for generating transgenic animals involves the genetic manipulation of embryonic stem cells (ES cells) as described in PCT Patent Publication No. WO 93/02188 and Robertson, 1987. In accordance with this technique, ES cells are grown as described in, for example, Robertson, 1987, and in U.S. Patent No. 5,166,065 to Williams et al. Genetic material is introduced into the embryonic stem cells by, for example, electroporation according, for example, to the method of McMahon, et al., 1990, or by transduction with a retroviral vector according, for example, to the method of Robertson, et al., 1986, or by any of the various techniques described by Lovell-Badge, 1987.

Chimeric animals are generated as described, for example, in Bradley, 1987. Briefly, genetically modified ES cells are introduced into blastocysts and the modified blastocysts are then implanted in pseudo-pregnant female animals. Chimeras are selected from the offspring, for example by the observation of mosaic coat coloration resulting from differences in the strain used to prepare the ES cells and the strain used to prepare the blastocysts, and are bred to produce non-chimeric transgenic animals.

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Other methods for the production of transgenic animals are disclosed in U.S. Patent No. 5,032,407 to Wagner et al., and PCT Publication No. WO90/08832.

Among other applications, transgenic prepared in accordance with the invention are useful as model systems for testing the xenotransplantation of their engineered tissues or organs and as sources of engineered tissues or organs for xenotransplantation. The expression of functional TMTCIPs on the surfaces of endothelial cells and/or other cell types in the tissues and organs (e.g., hormone producing cells such as those in the pancreatic islets) of the transgenic animals will provide enhanced protection to those cells, tissues and organs from hyperacute complement-mediated rejection following xenotransplantation in recipient animals, e.g., humans, whose complement can be inhibited by the Ly-6 terminal CIP of the TMTCIP. In addition to their use in producing organs for transplantation, the TMTCIP nucleic acid constructs of the invention can also be used to engineer cultured cells (e.g., endothelial cells) of various species for subsequent use in transplantation.

V. <u>Representative Modifications</u>

Although specific embodiments of the invention are described and illustrated herein, it is to be understood that modifications can be made without departing from the invention's spirit and scope.

For example, the primary amino acid structures of the TMTCIPs of the invention may be modified by creating amino acid substitutions or nucleic acid mutations. At least some complement regulatory activity should remain after such modifications. Similarly, nucleic acid mutations which do not change the amino acid sequences, e.g., third nucleotide changes in degenerate codons, are included within the scope of the invention. Also included are sequences comprising changes that are found as naturally occurring allelic variants of the CIP and TM genes used to create the TMTCIPs.

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Without intending to limit it in any manner, the present invention will be more fully described by the following examples.

Example 1

Expression Vectors and Retroviral Virion Particles Comprising a CD59/MCP TMTCIP

form of CD59 (CD59-TM) transmembrane constructed according to the present invention by replacing the carboxyl-terminal region containing the GPI- anchor signal of CD59 with the carboxyl-terminal region, including the transmembrane domain, An approximately 314 bp restriction fragment (CD46). (hereinafter referred to as CD5977) containing CD59 truncated at the "truncation-Asn" described above, i.e., amino acid 77 of the mature protein, was prepared by digestion of plasmid pCD59/CCPH (see below) with SspI and BamHI.

The carboxyl-terminus of CD46 was PCR amplified using HeLa cell reverse-transcribed mRNA as template and primers: 5'-CGCGAGGCCT ACTTACAAGC following (SEO ID NO:9) and 5'-CGCGCTATTC AGCCTCTCTG CTCCAG-3' ID NO:10). These oligonucleotides CTCTGC-3' (SEO amplified a fragment coding for amino acids 270-350 of the mature CD46 protein, a region shown previously to comprise a functional transmembrane domain (Lublin, et al., 1991). The approximately 250 bp fragment produced by this PCR reaction was cloned into a plasmid vector using the T/A cloning kit (Invitrogen, San Diego, CA). The pCRII plasmid vector included in this kit served as the recipient, and the resulting plasmid construct was amplified in E. coli and purified. The MCP insert was subsequently sequenced to confirm that the plasmid contained the sequence shown in SEQ ID NO:11.

An endogenous <u>Stu</u>I site found at the 5' end of the CD46 PCR fragment was utilized to ligate this domain to the <u>Ssp</u>I site at the 3' end of CD59₇₇ in the eukaryotic expression vector pcDNA3 (Invitrogen, San Diego, CA) to

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yield plasmid pcDNA3/CD59-MCP-TM (ATCC designation 69530).

The resulting construct was linearized with EcoRI, the unpaired ends were filled, and BamHI linkers (#1071, New England Biolabs, Tozer, MA) were ligated onto the resulting blunt ends. This linkered construct was digested with BamHI and the liberated fragment was subcloned into the BamHI site of the retroviral vector pLXSN (Miller, et al., 1989) to yield pL-CD59-MCP-TM-SN. Constructs with the correct orientation for expression were identified by restriction enzyme analysis and confirmed by sequencing.

A DNA fragment encoding the carboxyl-terminus of the CCPH gene was prepared by PCR amplification using plasmid pKS-/mCCPH (ATCC designation 69178) as template and the following primers: 5'-CCGGACCTGT GTAACTTTAA CGAACAGCTT GAAAATATTG GTAGGATATG CAATGGAAAT TGTTACAAC-3' (SEQ ID NO:12) and 5'-TAGTTACTGC CCGGACATGC-3' (SEQ ID NO:13). As described above for the MCP PCR fragment, the approximately 250 bp CCPH PCR product was cloned into plasmid pCRII, yielding plasmid pCRII/CCPH, and the CCPH insert was sequenced to confirm that the plasmid contained the desired sequence, in this case SEQ ID NO:14.

The pCRII/CCPH plasmid was then digested with <u>AvaII</u> and <u>EcoRI</u>, and the insert fragment was purified and subcloned in a three-way ligation reaction with plasmid pcDNA/AMP (Invitrogen) cut with <u>BamHI</u> and <u>EcoRI</u> and an approximately 300 base pair <u>BamHI</u> - <u>AvaII</u> fragment isolated from a full length CD59 cDNA construct in pUC19 (Philbrick et al., 1990). The product of this three-way ligation is referred to herein as plasmid pCD59/CCPH.

This plasmid was transfected into Balb/3T3 cells and the cells were assayed for cell surface expression of CD59 epitopes by indirect immunofluorescence as described below in Example 2. As discussed above, the putative TM domain of CCPH contains two cytoplasmic amino acids,

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within five amino acids of the first cytoplasmic amino acid, that are cysteine residues, a characteristic that is believed to result in low levels of cell surface expression. Cell surface expression of CD59 epitopes was indeed below the levels detectable by the indirect immunofluorescence assay.

Control vectors.

Full-length CD59 containing the GPI-anchor signal (CD59-GPI) was cloned into <u>BamHI</u> - <u>EcoRI</u> digested pcDNA3 (Invitrogen) as an <u>BamHI</u> - <u>EcoRI</u> fragment obtained from plasmid pc8-hCD59-103 (ATCC designation 69231) to yield plasmid pcDNA3-CD59-GPI.

Retroviral vector plasmid pL-CD59-GPI-SN was produced by isolating an approximately 1100 bp <u>EcoRI</u> fragment from a full length CD59 cDNA construct in pUC19 (Philbrick et al., 1990) and ligating this fragment into plasmid pLXSN. Constructs with the correct orientation for expression were identified by restriction enzyme analysis.

20 Amphotropic virus production.

was produced through Amphotropic virus intermediate ecotropic packaging cell line as described in Warren et al., 1987. Briefly, psi 2 cells (obtained from Dr. Stephen L. Warren, Department of Pathology, Yale University School of Medicine, New Haven, CT) were transfected with pLXSN or the pLXSN constructs described above, i.e., pL-CD59-MCP-TM-SN or pL-CD59-GPI-SN, using DMSO shock followed by selection in DMEM containing 500 (active) G418 and 10% heat inactivated FCS. Transfectants were pooled and a 24 hour supernatant was harvested from the cells at 90% confluency. The ecotropic virus stock was used to infect the amphotropic packaging cell line PA317 (ATCC designation CRL 9078). These cells were also selected in the same medium with G418 following which a virus stock was collected from pooled transductants in the same medium without G418.

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Example 2

Expression of the CD59/MCP TMTCIP by Mammalian Cells

Cells of the murine fibroblast cell line, Balb/3T3 (ATCC designation CCL 163) were stably transfected with pcDNA3-CD59-GPI, pcDNA3/CD59-MCP-TM, or pcDNA3 alone using the calcium phosphate method (Ausubel, et al., 1992). Cells were selected in DMEM containing 10% heat inactivated FCS and 500 μ g/ml of G418 (active) and colonies were isolated using cloning cylinders.

Mouse L cells were obtained from Dr. Immunobiology Department, Yale University Such mouse L cells School of Medicine, New Haven, CT. are unable to express GPI anchored proteins (Ferguson, et al., 1988). L cells were transduced with the amphotropic virus supernatants obtained using pL-CD59-GPI-SN. pL-CD59-MCP- TM-SN, or pLXSN alone by adding 1 ml of the virus stock to 5 x 10^5 L cells in medium containing 8 μg/ml polybrene. After an overnight incubation, medium containing 500 μ g/ml G418 was added and selection continued for 14 days. Transduced L cells were selected and analyzed as a pool. G418 resistant assayed for the presence of CD59 antiqens on the cell surface by indirect immunofluorescence using monoclonal and polyclonal antibody preparations. A rabbit polyclonal anti-CD59 antibody preparation, #349, that was produced by injecting rabbits with CD59 purified from human erythrocytes as described by Sims et al., 1989, was provided by Dr. Peter Sims (Blood Research Institute. Milwaukee, WI). The anti- CD59 mAb, MEM-43, purchased from Biodesign International, Kennebunkport, ME.

Cell surface indirect immunofluorescence analysis was typically performed on 2.5x10 5 cells with 50 $\mu g/ml$ of the primary polyclonal antibody or 20 $\mu g/ml$ of the monoclonal antibody in 1xPBS containing 2% fetal bovine serum. Goat anti-rabbit IgG or goat anti-mouse IgG FITC conjugated antisera were used as secondary antibodies

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(Zymed Laboratories, South San Francisco, CA). Fluorescence was measured using a FACSort instrument (Becton-Dickinson Immunocytometry Systems, San Jose, CA).

FIG. 2 illustrates cell surface expression profiles of a positive Balb/3T3 clone expressing CD59-TM, as well as of a native human CD59 (CD59-GPI) transfectant as a positive control and a vector (pcDNA3) without insert (vector control) transfectant as a negative control. As shown therein, essentially the same amount of anti-CD59 antibody bound to the surfaces of cells expressing the CD59-TM fusion protein as bound to the positive control cells expressing native CD59. This result shows that equivalent amounts of CD59 antigens were present on the Balb/3T3 cells of the invention (CD59-TM) and those of the positive control (CD59-GPI).

The pooled L cell transfectants showed a wide range of CD59-TM expression while, as expected in cells that cannot express GPI anchored proteins, CD59-GPI was not expressed (Figure 3).

Example 3

TMTCIP Expressed in Mammalian Cells Is Not Affected By Phosphatidylinositol-Phospholipase C Digestion

To test for the presence of a GPI anchor, cells were phosphatidylinositol-phospholipase treated with Boehringer-Mannheim Corporation, (PI-PLC, Products Division, Indianapolis, Indiana) at 1 U/ml for This treatment 1 hr at 37°C prior to FACS analysis. hydrolyzes (cleaves) GPI anchors, and thus frees GPI anchored proteins from the cell surface. digestion was performed on Balb/3T3 cells expressing the CD59-TM TMTCIP (or CD59-GPI as a control). The results of these experiments are presented in Figure 4. In these experiments, mock treated cells (no PI-PLC) retained the TMTCIP and native CD59 on their cell surfaces (see curve D in FIG. 4A and FIG. 4B), whereas PI-PLC treatment resulted in the loss of cell surface CIPs from the native CD59 control cells (see curve C in FIG. 4A), but not the CD59-TM cells (see curve C in FIG. 4B). These experiments demonstrate that CD59-TM is not anchored to the cell membrane through a GPI linkage and that CD59-TM is substantially resistant to the action of lipase enzymes which can cleave a glycosyl-phosphatidylinositol (GPI) anchor.

Example 4

Functional Analysis of CD59-TM in Mouse Cells

The functional activity of TMTCIP molecules expressed in transfected mouse Balb/3T3 cells and transduced mouse L cells was assessed by a dye release assay that consisted of measuring the efflux of molecules the cytoplasm, specifically the Calcein AM (Molecular Probes, indicator dye, Eugene, Oregon).

Transfected cells expressing the CD59-TM TMTCIP, as well as cells transfected with the parent expression vectors without CD59-TM encoding inserts (as controls), were grown to confluency in 96-well plates. Cells were washed twice with 200 μl of Hank's balanced salts solution containing 10 mg/ml bovine serum albumin (HBSS/BSA).

Calcein AM was added (10 μ M final) and the plates were incubated at 37°C for 30 minutes to allow the dye to be internalized by the cells and converted by cellular esterases into a polar fluorescent derivative that is retained inside undamaged cells. The wells were then washed twice with HBSS/BSA to remove dye remaining outside the cells. The cells were then incubated with anti- Balb/3T3 IgG (2 mg/ml in HBSS/BSA), which served as an activator of the classical complement pathway. After a 30 minute incubation at 23°C, unbound IgG was washed away.

The cells were then incubated at 37°C for 30 minutes in the presence of human C8 deficient serum supplemented with purified C8 and C9 to allow complement-mediated

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damage to occur. Human C8 depleted serum, as well as purified C8 and C9, were obtained from Quidel Corporation, San Diego, CA. The medium bathing the cells was then transferred to a clean 96-well plate for fluorescence measurement.

Under the conditions of this assay, the fluorescent polar derivative of Calcein AM is only released into the medium bathing the test cells if the integrity of the Therefore, the is compromised. membranes fluorescence of the Calcein AM released into the medium bathing the test cells versus that retained in the cells provides an indirect, but accurate measure of the level of complement- mediated damage sustained by the cells. Remaining cell- associated dye was determined from a 1% SDS lysate of the cells retained in the 96-well culture This allowed the calculation of percent dye release using the following formulas: Total = released total) \times 100. + retained, and, % release = (released Fluorescence was measured using a Millipore CYTOFLUOR 2350 fluorescence plate reader (490 nm excitation, 530 nm emission).

The dye release assays showed that for transfected Balb/3T3 clones expressing equivalent levels of CD59-GPI or CD59-TM (FIG. 2), CD59-TM provided a level of protection from complement attack equivalent to that afforded by the native, GPI-anchored, CD59-GPI molecule (FIG. 5). In particular, cells expressing either of these molecules were approximately 3-fold more effective in preventing complement-mediated lysis at 2.5 μ g/ml C8/C9 than cells transfected with the pcDNA3 vector alone, which were readily lysed.

These results demonstrate that 1) CD59-TM can be stably expressed on the surface of Balb/3T3 cells, and 2) this chimeric molecule has comparable function to native CD59. The retention of wild-type levels of complement regulatory activity by CD59-TM is of considerable significance in that it shows that the functionality of

PCT/US95/02944

the CD59 molecule is not substantially altered by truncation coupled with addition of a TM domain. This result could not have been predicted in advance, especially since other alterations of the CD59 molecule, e.g., truncation of the carboxyl-terminus without addition of a TM domain, or alterations of single amino acids, have been shown to produce molecules with substantially altered expression and/or functionality. See, for example, Nakano, et al., 1993; Norris, et al., 1993; and Petranka, et al., 1993.

Dye release assays were also performed on mouse L cells transduced with the retroviral virion particles using the pL-CD59-MCP-TM-SN generated vector, pL-CD59- GPI-SN vector, or the no insert pLXSN vector. The results of these experiments are presented in FIG. 6. Only L cells transduced with retroviral particles generated using pL-CD59-MCP-TM-SN demonstrated substantial protection against complement attack. results demonstrate that the chimeric CD59-TM molecule can successfully be expressed in a cell line unable to express GPI-anchored proteins and that the molecule functions to protect the cells from complement lysis.

The foregoing results show that CD59 retains its Ly-6 terminal complement inhibitor activity when anchored to the cell membrane by a heterologous transmembrane domain, rather than a GPI anchor. This fundamental result, in combination with the conserved nature of all known Ly-6 terminal complement inhibitor proteins (see U.S. patent application Serial No. 08/105,735 and PCT patent application Serial No. PCT/US93/006772), indicates heterologous transmembrane domain a substituted for the GPI signal sequence of a Ly-6 terminal complement inhibitor protein substantially altering the complement inhibitor activity of the protein.

In comparison to using a native Ly-6 terminal CIP, the TMTCIPs of the invention have the advantages that

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they cannot produce cell activation of the type which depends on the presence of the GPI anchor of the native Ly-6 terminal CIP, and that they cannot be removed from the cell surface by the action of phospholipase enzymes and are less prone to vesicular shedding. These advantages make the TMTCIPs of the invention more suitable than native Ly-6 terminal CIPs for various medical applications, including the facilitation of transplantation of xenogenetic organs.

Although preferred and other embodiments of the invention have been described herein, other embodiments, including a variety of modifications may be perceived and practiced by those skilled in the art without departing from the scope of the invention. For example, the primary amino acid structures of the fusion proteins of the invention may be modified by creating amino acid mutants. Such mutants should retain more than 50% of the complement regulatory activity of the parent terminal CIP. Other modifications and variations include forming derivatives of the fusion protein to include covalent or aggregated conjugates of the protein or its fragments with other proteins or polypeptides. The following claims are intended to cover the specific embodiments set forth herein as well as such modifications, variations, and equivalents.

Throughout this application, various publications, patents, and patent applications have been referred to. The teachings and disclosures of these publications, patents, and patent applications in their entireties are hereby incorporated by reference into this application to more fully describe the state of the art to which the present invention pertains.

DEPOSITS

Plasmids pcDNA3/CD59-MCP-TM, pc8-hCD59-103, and pKS-/mCCPH, discussed above, have been deposited with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland, 20852, United States of America, in

<u>E. coli</u> and have been assigned the designations 69530, 69231, and 69178, respectively. These deposits were made under the Budapest Treaty on the International Recognition of the Deposit of Micro-organisms for the Purposes of Patent Procedure (1977).

The deposits referred to above having ATCC accession numbers 69530, 69231, and 69178 were made on January 6, January 29, 1993, and January 6, respectively. Deposit 69530 was made in Escherichia coli strain TOP10F' which has the following $F'\{lacI^q TN10(Tet^R)\} mcrA \Delta(mrr-hsdRMS-mcrBC) \phi80lacZ\DeltaM15$ ΔlacX74 deoR recA1 araD139 Δ(ara-leu)7697 galU galK rpsL(StrR) endA1 nupG. Deposits 69231 and 69178 were made in Escherichia coli strain DH5 α which has the following genotype: F φ80dlacZΔM15 Δ(lacZYA-argF)U169 recAl endAl $hsdR17(r_k-,m_k+)$ supE44 λ thi-1 gyrA96 relA1.

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT: Rother, Russell Rollins, Scott Squinto, Stephen P
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- (iii) NUMBER OF SEQUENCES: 14
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 - (C) CITY: Fairfield
 - (D) STATE: Connecticut
 - (E) COUNTRY: USA
 - (F) ZIP: 06430
- (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: 3.5 inch, 720 Kb storage
 - (B) COMPUTER: Dell 486/50
 - (C) OPERATING SYSTEM: DOS 6.2
 - (D) SOFTWARE: Word Perfect 6.0
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: 08/205,720
 - (B) FILING DATE: 3-MAR-1994
 - (C) CLASSIFICATION:

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(2)	TNFORMATTON	TOD	CEC	TD	NTO - 7 -

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 763 base pairs
 - (B) TYPE: Nucleic Acid
 - (C) STRANDEDNESS: Double
 - TOPOLOGY: Linear (D)
- (ii) MOLECULE TYPE: cDNA to mRNA
 - DESCRIPTION: BABCIP full length cDNA (A)
- (iii) HYPOTHETICAL: No
- (iv) ANTI-SENSE: No
- (vi) ORIGINAL SOURCE:
 - ORGANISM: Papio hamadryas (A)
- IMMEDIATE SOURCE: (vii)
 - (A) LIBRARY: Baboon Spleen Lambda ZAPII cDNA Library, Catalog # 936103, Stratagene Cloning Systems, La Jolla, California
- SEQUENCE DESCRIPTION: SEQ ID NO:1:

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ATC CAA GGA GGG TCT GTC CTG TTC GGG CTG CTG CTT GTC CTG GCT 185 Ile Gln Gly Gly Ser Val Leu Phe Gly Leu Leu Leu Val Leu Ala -20 -15

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					GCG Ala									ACC Thr	365
					GAG Glu									GAC Asp	410
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GCAC	GGA <i>I</i>	AGC (CCA	CTTG	AA GO	GAAG	AAGT:	TAI	AGAG:	rgaa	GTA	GTG	rga		748
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(C) STRANDEDNESS: Double
(D) TOPOLOGY: Linear
(ii) MOLECULE TYPE: cDNA to mRNA
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(iii) HYPOTHETICAL: No
(iv) ANTI-SENSE: No
(vi) ORIGINAL SOURCE:
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(H) CELL LINE: COS-1 (ATCC CRL 1650)
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:
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Gln Gly Ser Val Leu Phe Gly Leu Leu Ala Leu Ala Val -20 -15 -10
TTC TGC CAT TCA GGT CAT AGC CTG CAA TGC TAC AAC TGT CCT AAC 116 Phe Cys His Ser Gly His Ser Leu Gln Cys Tyr Asn Cys Pro Asn
1 -5 5 T
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GAT ACG TGT CTC ATT GCC AGA GCT GGG TTA CAA GTA TAT AAC CAG 206 Asp Thr Cys Leu Ile Ala Arg Ala Gly Leu Gln Val Tyr Asn Gln 25 30 35

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 - (B) TYPE: Nucleic Acid
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 - (D) TOPOLOGY: Linear
 - (ii) MOLECULE TYPE: cDNA to mRNA
 - (A) DESCRIPTION: SQMCIP full coding cDNA
 - (iii) HYPOTHETICAL: No
 - (iv) ANTI-SENSE: No
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Saimiri sciureus
 - (H) CELL LINE: DPSO 114/74 (ATCC CCL 194)
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:
- ATG GGA ATC CAA GGA GGG TCT GTC CTG TTT GGG CTG CTC GTC 45
 Met Gly Ile Gln Gly Gly Ser Val Leu Phe Gly Leu Leu Val
 -25
 -20
 -15
- CTG GCT GTC TGC CAT TCA GGT AAT AGC CTG CAA TGC TAC AGC 90 Leu Ala Val Phe Cys His Ser Gly Asn Ser Leu Gln Cys Tyr Ser -10 -5 1
- TGT CCT CTC CCA ACC ATG GAG TCC ATG GAG TGC ACT GCG TCC ACC 135 Cys Pro Leu Pro Thr Met Glu Ser Met Glu Cys Thr Ala Ser Thr 10 15 20
- AAC TGT ACA TCT AAT CTT GAT TCG TGT CTC ATT GCC AAA GCC GGG 180 Asn Cys Thr Ser Asn Leu Asp Ser Cys Leu Ile Ala Lys Ala Gly 25 30 35
- TCA GGA GTA TAT TAC CGG TGT TGG AAG TTT GAC GAT TGC AGT TTC 225 Ser Gly Val Tyr Tyr Arg Cys Trp Lys Phe Asp Asp Cys Ser Phe 40 45 50
- AAA CGC ATC TCA AAC CAA TTG TCG GAA ACT CAG TTA AAG TAT CAC 270 Lys Arg Ile Ser Asn Gln Leu Ser Glu Thr Gln Leu Lys Tyr His 55 60 65

WO 95/23512 PCT/US95/02944

-55-

				GTT Val				AAT Asn 80	315
				ACA Thr				ACC Thr 95	360
				CGT Arg		TAA	٠	·	396

-56-

- INFORMATION FOR SEQ ID NO:4: (2)
 - SEQUENCE CHARACTERISTICS: (i)
 - LENGTH: 387 base pairs (A)
 - TYPE: Nucleic Acid (B)
 - STRANDEDNESS: Double (C)
 - (D) TOPOLOGY: Linear
 - MOLECULE TYPE: cDNA to mRNA (ii)
 - DESCRIPTION: OWMCIP full coding cDNA (A)
 - (iii) HYPOTHETICAL: No
 - ANTI-SENSE: No (iv)
 - ORIGINAL SOURCE: (vi)
 - ORGANISM: Aotus trivirgatus (A)
 - CELL LINE: OMK (ATCC CRL 1556) (H)
 - (xi) SEOUENCE DESCRIPTION: SEQ ID NO:4:
- ATG GGA ATT CAA GGA GGG TCT GTC CTG TTT GGG CTG CTC GTC Met Gly Ile Gln Gly Gly Ser Val Leu Phe Gly Leu Leu Val
- CTG GCT GTC TTC TGC CAT TCA GGT AAT AGC CTG CAG TGC TAC AGC Leu Ala Val Phe Cys His Ser Gly Asn Ser Leu Gln Cys Tyr Ser -10
- TGT CCT TAC CCA ACC ACT CAG TGC ACT ATG ACC ACC AAC TGT ACA 135 Cys Pro Tyr Pro Thr Thr Gln Cys Thr Met Thr Thr Asn Cys Thr
- TCT AAT CTT GAT TCG TGT CTC ATT GCC AAA GCC GGG TCA CGA GTA 180 Ser Asn Leu Asp Ser Cys Leu Ile Ala Lys Ala Gly Ser Arg Val
- TAT TAC CGG TGT TGG AAG TTT GAG GAT TGC ACT TTC AGC CGC GTT 225 Tyr Tyr Arg Cys Trp Lys Phe Glu Asp Cys Thr Phe Ser Arg Val
- TCA AAC CAA TTG TCT GAA AAT GAG TTA AAG TAT TAC TGC TGC AAG 270 Ser Asn Gln Leu Ser Glu Asn Glu Leu Lys Tyr Tyr Cys Cys Lys 55 60

-57-

		AAC Asn 70						315
		AAA Lys 85						360
		AGC Ser 100		TAA				387

-58-

INFORMATION FOR SEO ID NO:5: (2)

- SEQUENCE CHARACTERISTICS: (i)
 - LENGTH: 387 base pairs (A)
 - (B) TYPE: Nucleic Acid
 - STRANDEDNESS: Double (C)
 - TOPOLOGY: Linear (D)
- MOLECULE TYPE: cDNA to mRNA (ii)
 - DESCRIPTION: MARCIP full coding cDNA (A)
- (iii) HYPOTHETICAL: No
- ANTI-SENSE: No (iv)
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Saguinus nigricollis
 - (H) CELL LINE: 1283.Lu (ATCC CRL 6297)
- SEQUENCE DESCRIPTION: SEQ ID NO:5:
- ATG GGA ATC CAA GGA GGG TCT GTC CTG TTT GGG CTG CTC ATC Met Gly Ile Gln Gly Gly Ser Val Leu Phe Gly Leu Leu Ile
- CTG GCT GTC TTC TGC CAT TCA GGT CAT AGC CTG CAG TGC TAC AGC Leu Ala Val Phe Cys His Ser Gly His Ser Leu Gln Cys Tyr Ser
- TGT CCT TAC TCA ACC GCT CGG TGC ACT ACG ACC ACC AAC TGT ACA 135 Cys Pro Tyr Ser Thr Ala Arg Cys Thr Thr Thr Thr Asn Cys Thr 10 15
- TCT AAT CTT GAT TCA TGT CTC ATT GCC AAA GCC GGG TTA CGA GTA 180 Ser Asn Leu Asp Ser Cys Leu Ile Ala Lys Ala Gly Leu Arg Val
- TAT TAC CGG TGT TGG AAG TTT GAG GAT TGC ACT TTC AGA CAA CTT 225 Tyr Tyr Arg Cys Trp Lys Phe Glu Asp Cys Thr Phe Arg Gln Leu
- TCA AAC CAA TTG TCG GAA AAT GAG TTA AAG TAT CAC TGC TGC AGG 270 Ser Asn Gln Leu Ser Glu Asn Glu Leu Lys Tyr His Cys Cys Arg 55 60

	AAC Asn													ACA Thr 80	315
ACC Thr	TTA Leu	TCA Ser	AAG Lys	AAA Lys 85	ACA Thr	GTT Val	CTT Leu	CTG Leu	CTG Leu 90	GTG Val	ACC Thr	CCT Pro	TTT Phe	CTG Leu 95	360
	GCA Ala							TAA							387

i

(2) INFORMATION FOR	SEO	ID	NO:6:
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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1039 base pairs
 - (B) TYPE: Nucleic Acid
 - (C) STRANDEDNESS: Double
 - (D) TOPOLOGY: Linear
- (ii) MOLECULE TYPE: cDNA to mRNA
 - (A) DESCRIPTION: HVS-15 full length cDNA
- (iii) HYPOTHETICAL: No
- (iv) ANTI-SENSE: No
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Herpesvirus saimiri
- (x) PUBLICATION INFORMATION:
 - (A) AUTHORS: Albrecht, J.C.
 Nicholas, J.
 Cameron. K.R.
 Newman, C.
 Fleckenstein, B.

Honess, R.W.

(B) TITLE: Herpesvirus samiri has a gene specifying a homologue of the cellular membrane

glycoprotein CD59.

- (C) JOURNAL: Virology
- (D) VOLUME: 190
- (F) PAGES: 527-530
- (G) DATE: 1992

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

AAGCTTCTAT TTATACTACA TTAGAGGCAT TTTTTCAAAA GCAAAAATGC 50
CTCTAATTAT ATACACTGTA CTATTTACCT CTATTACACA TTTTCTATTT 100
TAAGTCTGAT AGTGATTAAT CAAGAAAAAA GTTTGTGGTT CTCAGGGGAT 150
TAGTTCACAA GCTGTCTGAG GTTAAGGGTG TTTCTTTGGC ACTGACACAG 200
AAGTTGCTAT AAGAATTGAA GCTTGCTTTA CAAAAAGTTA CTTGTGATTA 250

ATT	ACTA!	CAT	CAAGI	AAAG	GT A						ACG Thr			295
CTG Leu	ACT Thr -10	TTT Phe	GTT Val	TTT	TGC Cys	AAG Lys -5	CCA Pro	ATA Ile	CAC His	AGC Ser	TTG Leu 1	CAA Gln	TGC Cys	337
TAC Tyr	AAC Asn 5	TGT Cys	TCT Ser	CAC His	TCA Ser	ACT Thr 10	ATG Met	CAG Gln	TGT Cys	ACT Thr	ACA Thr 15	TCT Ser	ACT Thr	379
AGT Ser	TGT Cys	ACA Thr 20	TCT Ser	AAT Asn	CTT Leu	GAC Asp	TCT Ser 25	TGT Cys	CTC Leu	ATT Ile	GCT Ala	AAA Lys 30	GCT Ala	421
GGG Gly	TCA Ser	GGA Gly	GTA Val 35	TAT Tyr	TAC Tyr	AGG Arg	TGT Cys	TGG Trp 40	AAG Lys	TTT Phe	GAT Asp	GAC Asp	TGT Cys 45	463
AGC Ser	TTT Phe	AAA Lys	CGT Arg	ATC Ile 50	TCA Ser	AAT Asn	CAA Gln	TTG Leu	TCT Ser 55	GAA Glu	ACA Thr	CAG Gln	TTA Leu	505
AAG Lys 60	TAT Tyr	CAT His	TGT Cys	TGT Cys	AAG Lys 65	AAG Lys	AAC Asn	TTG Leu	TGT Cys	AAT Asn 70	GTG Val	AAC Asn	AAA Lys	547
GGG Gly	ATT Ile 75	GAA Glu	AAT Asn	ATŢ Ile	AAA Lys	AGA Arg 80	ACA Thr	ATA Ile	TCA Ser	GAT Asp	AAA Lys 85	GCT Ala	CTT Leu	589
TTA Leu	CTA Leu	TTA Leu 90	GCA Ala	TTG Leu	TTT Phe	TTA Leu	GTA Val 95	ACT Thr	GCT Ala	TGG Trp	AAC Asn	TTT Phe 100	CCT Pro	631
CTT Leu	TAAA	AAG :	CAAC	CAACA	AA AA	CTAT	TATTO	TAI	CATT	TAT	TTT]	rgtgt	rag	680
CTT	TTTC	TA T	rtgci	TATTA	AC AA	GTT	CAAA	r ATT	GTGT	TTT	TTA	ACTA	ATA	730
ATT	ATT	AAA	AGATZ	CAAA!	rg ac	ATGI	TAGT!	A TAC	CTACA	ATAG	TCA	TAAL	AAT	780
AGTO	CTA	AT A	TATTA	TAGO	CA AT	TTTT	TATO	AAC	CAACO	CAA	ATA	AAGI	TA	830
AGC	ract"	TA :	rtttt	TCTC	T T	ATCTZ	TAAL	CATI	ACG	CGCT	TCTT	TAGC	ATG	880
TGTT	\AAA'	AGT T	ratta	GTGI	TT T	TATT	CTTA	A CAI	CATA	AAA	GCT	LTAAL	TT	930
AAA	CAA	ATT A	ATCAG	TAGO	CA TO	TTAT	CTTC	TAF	ATCTO	TAC	AGAC	CTAT	TAT	980
AATZ	ATGGG	TAS	TATCO	TTA	AG AA	LAAA	ACAGO	C GGA	GAAZ	AAAG	AAAA	ACACA	AGT	1030
GCC	AAGCI	ГT												1039

- (2) INFORMATION FOR SEQ ID NO:7:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1139 base pairs
 - (B) TYPE: Nucleic Acid
 - (C) STRANDEDNESS: Double
 - (D) TOPOLOGY: Linear
 - (ii) MOLECULE TYPE: cDNA to mRNA
 - (A) DESCRIPTION: CD59 full length cDNA
 - (iii) HYPOTHETICAL: No
 - (iv) ANTI-SENSE: No
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo sapiens
 - (x) PUBLICATION INFORMATION:
 - (A) AUTHORS: Philbrick, W.M.
 Palfree, R.G.E
 Maher, S.E.
 Bridgett, M.M.
 Sirlin S.

Bothwell, A.L.M.

- (B) TITLE: The CD59 antigen is a structural homologue of murine Ly-6 antigens but lacks interferon inducibility.
- (C) JOURNAL: European Journal of Immunology
- (D) VOLUME: 20
- (F) PAGES: 87-92
- (G) DATE: JAN-1990

-63-

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

CGCAGAAGCG GCTCGAGGCT GGAAGAGGAT CCTGGGCGCC GCAGGTTCTG										50				
	TGGAC	ATCA	M	TG G(et G: 25				ly G						92
	GGG CT Gly Le -15	rg CTo	G CTC u Leu	GTC Val	CTG Leu -10	GCT Ala	GTC Val	TTC Phe	TGC Cys	CAT His	TCA Ser	GGT Gly	CAT His	134
	AGC CT Ser Le	rg CA eu Gla L	G TGC n Cys	TAC Tyr	AAC Asn 5	TGT Cys	CCT Pro	AAC Asn	CCA Pro	ACT Thr 10	GCT Ala	GAC Asp	TGC Cys	176
	AAA AC Lys Th	CA GCO ar Ala	C GTC a Val	AAT Asn	TGT Cys	TCA Ser 20	TCT Ser	GAT Asp	TTT Phe	GAT Asp	GCG Ala 25	TGT Cys	CTC Leu	218
	ATT AC	CC AAI	s Ala	GGG Gly	TTA Leu	CAA Gln	GTG Val 35	TAT Tyr	AAC Asn	AAG Lys	TGT Cys	TGG Trp 40	AAG Lys	260
	TTT GAP Phe Gl	AG CA' Lu His	TGC Cys 45	AAT Asn	TTC Phe	AAC Asn	GAC Asp	GTC Val 50	ACA Thr	ACC Thr	CGC Arg	TTG Leu	AGG Arg 55	302
	GAA AA Glu As	AT GAG	G CTA 1 Leu	ACG Thr 60	TAC Tyr	TAC Tyr	TGC Cys	TGC Cys	AAG Lys 65	AAG Lys	GAC Asp	CTG Leu	TGT Cys	344
	AAC TT Asn Ph 70	TT AAG	C GAA n Glu	CAG Gln	CTT Leu 75	GAA Glu	AAT Asn	GGT Gly	GGG Gly	ACA Thr 80	TCC Ser	TTA Leu	TCA Ser	386
	GAG AA Glu Ly	AA AC 7s Th: 35	A GTT r Val	CTT Leu	CTG Leu	CTG Leu 90	GTG Val	ACT Thr	CCA Pro	TTT Phe	CTG Leu 95	GCA Ala	GCA Ala	428
	GCC TO		r Leu			TAA	G T	CAAC	ACCA(G GA	GAGC:	TCT		470
	CCCAA	CTCC	CCGT"	TCCT	GC G	ragt(CCGC'	r TT	CTCT	rgct	GCC	ACAT"	rct	520
	AAAGG	CTTGA	TATT'	TTCC	AA A	rgga:	rcct(G TTC	EGGA.	AAGA	ATA	TAAL	ràg	570
	CTTGAC	CAAC	CTGG	CTAA	GA T	AGAG	GGT	C TG	GAG	ACTT	TGA	AGAC	CAG	620
	TCCTG	CCCGC	AGGG.	AAGC	cc cz	ACTT	GAAG	G AA	GAAG'	rcta	AGA	GTGA/	AGT	670
	AGGTGT	rgact	TGAA	CTAG	AT TO	GCAT(GCTT(CT	CCTT	TGCT	CTT	GGA	AGA	720

PCT/US95/02944

WO 95/23512

-64-

CCAGCTTTGC	AGTGACAGCT	TGAGTGGGTT	CTCTGCAGCC	CTCAGATTAT	770
TTTTCCTCTG	GCTCCTTGGA	TGTAGTCAGT	TAGCATCATT	AGTACATCTT	820
TGGAGGGTGG	GGCAGGAGTA	TATGAGCATC	CTCTCTCACA	TGGAACGCTT	870
TCATAAACTT	CAGGGATCCC	GTGTTGCCAT	GGAGGCATGC	CAAATGTTCC	920
ATATGTGGGT	GTCAGTCAGG	GACAACAAGA	TCCTTAATGC	AGAGCTAGAG	970
GACTTCTGGC	AGGGAAGTGG	GGAAGTGTTC	CAGATTCCAG	ATAGCAGGGC	1020
ATGAAAACTT	AGAGAGGTAC	AAGTGGCTGA	AAATCGAGTT	TTTCCTCTGT	1070
CTTTAAATTT	TATATGGGCT	TTGTTATCTT	CCACTGGAAA	AGTGTAATAG	1120
CATACATCAA	TGGTGTGTT				1139

-65-

INFORMATION FOR SEQ ID NO:8: (2)

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1530 base pairs
 - (B) TYPE: Nucleic Acid
 - (C) STRANDEDNESS: Double
 - (D) TOPOLOGY: Linear
- (ii) MOLECULE TYPE: cDNA to mRNA
 - (A) DESCRIPTION: MCP (CD46) full length cDNA
- (iii) HYPOTHETICAL: No
- (iv) ANTI-SENSE: No
- (vi) ORIGINAL SOURCE:
 - . (A) ORGANISM: Homo sapiens
- PUBLICATION INFORMATION: (x)
 - (A) **AUTHORS:** Lublin, D.M. Liszewski, M.K. Post, T.W. Arce, M.A. LeBeau, M.M. Rebentisch, M.B. Lemons, R.S. Seya, T.

Atkinson, J.P.

(B) TITLE: Molecular cloning and Chromosomal Localization of Membrane Cofactor

Protein (MCP): Evidence for

Inclusion in the Multi-Gene Family of Complement-Regulatory Proteins.

- (C) JOURNAL: Journal of Experimental Medicine
- (D) VOLUME: 168
- (F) PAGES: 181-194
- (G) DATE: 1988

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

TCTGCTTTCC	TCCGGAGA	AA TAACA	GCGTC T	rccgcgccg	CGC	ATG Met		49
CCT CCC GG Pro Pro Gl -3	/ Arg Arg							91
CCT GGG TT Pro Gly Le	G CTT CTG Leu Leu -15	GCG GCC Ala Ala	ATG GTO Met Val	Leu Leu	CTG Leu	TAC Tyr	TCC Ser -5	133
TTC TCC GA Phe Ser As	GCC TGT Ala Cys 1	GAG GAG Glu Glu	CCA CCA Pro Pro	A ACA TTT Thr Phe	GAA Glu	GCT Ala	ATG Met 10	17 5
GAG CTC AT Glu Leu Il	GGT AAA Gly Lys 15	CCA AAA Pro Lys	CCC TAC Pro Ty	TAT GAG Tyr Glu 20	ATT Ile	GGT Gly	GAA Glu	217
CGA GTA GA Arg Val As 25	TAT AAG Tyr Lys	TGT AAA Cys Lys 30	AAA GGA	TAC TTC Tyr Phe 35	TAT Tyr	ATA Ile	CCT Pro	259
CCT CTT GC Pro Leu Al 40	C ACC CAT A Thr His	ACT ATT Thr Ile 45	Cys Asp	CGG AAT Arg Asn	CAT His 50	ACA Thr	TGG Trp	301
CTA CCT GT Leu Pro Va 5	L Ser Asp	GAC GCC Asp Ala	TGT TAT Cys Tyr 60	T AGA GAA T Arg Glu	ACA Thr	TGT Cys 65	CCA Pro	343
TAT ATA CG Tyr Ile Ar	GAT CCT Asp Pro 70	TTA AAT Leu Asn	GGC CAA Gly Glr 75	n Ala Val	CCT Pro	GCA Ala	AAT Asn 80	385
GGG ACT TA Gly Thr Ty	GAG TTT Glu Phe 85	GGT TAT Gly Tyr	CAG ATO	CAC TTT His Phe 90	ATT Ile	TGT Cys	AAT Asn	427
GAG GGT TA Glu Gly Ty 95	TAC TTA Tyr Leu	Ile Gly	GAA GAA Glu Glu	A ATT CTA 1 Ile Leu 105	TAT Tyr	TGT Cys	GAA Glu	469
CTT AAA GG Leu Lys Gl 110	A TCA GTA V Ser Val	GCA ATT Ala Ile 115	Trp Ser	C GGT AAG Gly Lys	CCC Pro 120	CCA Pro	ATA Ile	511
TGT GAA AA Cys Glu Ly 12	s Val Leu	TGT ACA Cys Thr	CCA CCT Pro Pro 130	CCA AAA Pro Lys	ATA Ile	AAA Lys 135	AAT Asn	553

GGA A														595
GAT G Asp A														637
CCA T Pro P 165														679
AAT I Asn S														721
AAA I	Cys	CGA Arg 195	TTT Phe	CCA Pro	GTA Val	GTC Val	GAA Glu 200	AAT Asn	GGA Gly	AAA Lys	CAG Gln	ATA Ile 205	Ser	763
GGA T	TTT Phe	GGA Gly	AAA Lys 210	AAA Lys	TTT Phe	TAC Tyr	TAC Tyr	AAA Lys 215	Ala	ACA Thr	GTT Val	ATG Met	TTT Phe 220	805
GAA T Glu C	rgc Cys	GAT Asp	AAG Lys	GGT Gly 225	TTT Phe	TAC Tyr	CTC Leu	GAT Asp	GGC Gly 230	AGC Ser	GAC Asp	ACA Thr	ATT Ile	847
GTC T Val C 235	rgt Cys	GÄC Asp	AGT Ser	AAC Asn	AGT Ser 240	ACT Thr	TGG Trp	GAT Asp	CCC Pro	CCA Pro 245	GTT Val	CCA Pro	AAG Lys	889
TGT C Cys L 2	Leu 250	AAA Lys	GTG Val	TCG Ser	ACT Thr	TCT Ser 255	TCC Ser	ACT Thr	ACA Thr	AAA Lys	TCT Ser 260	CCA Pro	GCG Ala	931
TCC A Ser S	Ser	GCC Ala 265	TCA Ser	GGT Gly	CCT Pro	AGG Arg	CCT Pro 270	ACT Thr	TAC Tyr	AAG Lys	CCT Pro	CCA Pro 275	GTC Val	973
TCA A							Lys							1015
GAC A Asp S	AGT Ser	TTG Leu	GAT Asp	GTT Val 295	TGG Trp	GTC Val	ATT Ile	GCT Ala	GTG Val 300	ATT Ile	GTT Val	ATT Ile	GCC Ala	1057
ATA G Ile V 305	TT /al	GTT Val	GGA Gly	GTT Val	GCA Ala 310	GTA Val	ATT Ile	TGT Cys	GTT Val	GTC Val 315	CCG Pro	TAC Tyr	AGA Arg	1099
TAT C Tyr I	CTT Leu 320	CAA Gln	AGG Arg	AGG Arg	AAG Lys	AAG Lys 325	AAA Lys	GGG Gly	AAA Lys	GCA Ala	GAT Asp 330	GGT Gly	.GGA Gly	1141

	Ala Thr Ty			ACT CCA GCA Thr Pro Ala 345	1183
GAG CAG AGA Glu Gln Arg		F AGATTCCACA	A ACCTGGTTTG	CCAGTTCATC	1230
TTTTGACTCT	ATTAAAATCT	TCAATAGTTG	TTATTCTGTA	GTTTCACTCT	1280
CATGAGTGCA	ACTGTGGCTT	AGCTAATATT	GCAATGTGGC	TTGAATGTAG	1330
GTAGCATCCT	TTGATGCTTC	TTTGAAACTT	GTATGAATTT	GGGTATGAAC	1380
AGATTGCCTG	CTTTCCCTTA	AATAACACTT	AGATTTATTG	GACCAGTCAG	1430
CACAGCATGC	CTGGTTGTAT	TAAAGCAGGG	ATATGCTGTA	TTTTATAAAA	1480
ייירככר א א א א יי	тасасааата	ጥ ልርጥጥርልሮልል	יים אם אריד מיד מ	սուրարարարա	1530

PCT/US95/02944

-69-

(2)	INFORMATION	FOR	SEO	TD	NO:9:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 26 bases
 - (B) TYPE: Nucleic Acid
 - (C) STRANDEDNESS: Single
 - (D) TOPOLOGY: Linear
- (ii) MOLECULE TYPE: Other nucleic acid
- (iii) HYPOTHETICAL: No
- (iv) ANTI-SENSE: No
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

CGCGAGGCCT ACTTACAAGC CTCCAG

26

- (2) INFORMATION FOR SEQ ID NO 10:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 26 bases
 - (B) TYPE: Nucleic Acid
 - (C) STRANDEDNESS: Single
 - (D) TOPOLOGY: Linear
 - (ii) MOLECULE TYPE: Other nucleic acid
 - (iii) HYPOTHETICAL: No
 - (iv) ANTI-SENSE: Yes
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

CGCGCTATTC AGCCTCTCTG CTCTGC

TGAATAGCGC G

(2) INFORMATION FOR SEQ ID NO:11:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 261 base pairs	
(B) TYPE: Nucleic Acid	
(C) STRANDEDNESS: Double	
(D) TOPOLOGY: Linear	
(ii) MOLECULE TYPE: Other nucleic acid	
(A) DESCRIPTION: MCP PCR Product	
(iii) HYPOTHETICAL: No	
(iv) ANTI-SENSE: No	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:	
CGCGAGGCCT ACTTACAAGC CTCCAGTCTC AAATTATCCA GGATATCCTA	50
AACCTGAGGA AGGAATACTT GACAGTTTGG ATGTTTGGGT CATTGCTGTG	100
ATTGTTATTG CCATAGTTGT TGGAGTTGCA GTAATTTGTG TTGTCCCGTA	150
CAGATATCTT CAAAGGAGGA AGAAGAAAGG GAAAGCAGAT GGTGGAGCTG	200
AATATGCCAC TTACCAGACT AAATCAACCA CTCCAGCAGA GCAGAGAGGC	250

(2) INFORMATION FOR SEQ ID NO:12:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 69 bases	
(B) TYPE: Nucleic Acid	
(C) STRANDEDNESS: Single	
(D) TOPOLOGY: Linear	
(ii) MOLECULE TYPE: Other nucleic acid	
(iii) HYPOTHETICAL: No	
(iv) ANTI-SENSE: No	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:	
CCGGACCTGT GTAACTTTAA CGAACAGCTT GAAAATATTG GTAGGATATG	50
CAATGGAAAT TGTTACAAC	69
(2) INFORMATION FOR SEQ ID NO:13:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 20 bases	
(B) TYPE: Nucleic Acid	
(C) STRANDEDNESS: Single	
(D) TOPOLOGY: Linear	
(ii) MOLECULE TYPE: Other nucleic acid	
(iii) HYPOTHETICAL: No	
(iv) ANTI-SENSE: Yes	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
TAGTTACTGC CCGGACATGC	20

264

CCGGGCAGTA ACTA

(2) INFORMATION FOR SEQ ID NO:14:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 264 base pairs	
(B) TYPE: Nucleic Acid	
(C) STRANDEDNESS: Double	
(D) TOPOLOGY: Linear	
(ii) MOLECULE TYPE: Other nucleic acid	
(A) DESCRIPTION: CCPH PCR Product	
(iii) HYPOTHETICAL: No	
(iv) ANTI-SENSE: No	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	
CCGGACCTGT GTAACTTTAA CGAACAGCTT GAAAATATTG GTAGGATATG	50
CAATGGAAAT TGTACAACTA GCATGCCCAC TCAAACATAT ACAATAATTA 1	00
CTGCGCGCTA TACAAGTCAC ATATATTTCC CTACTGGGAA AACCTATAAA 1	50
CTTCCTCGGG GAGTTCTAGT AATTATTCTT ACCACAAGCT TTATTATTAT 2	00
TGGAATAATA CTTACTGGAG TGTGTTTACA TAGGTGCAGA GTGTGCATGT 2	50

WO 95/23512 PCT/US95/02944

What is claimed is:

- A nucleic acid molecule comprising:
- a sequence encoding a chimeric protein which comprises:

-73-

- (i) a first polypeptide region comprising a portion of a parent terminal complement inhibitor protein, said portion including a complete Ly-6 motif and not including an operative signal sequence directing the attachment of a glycosyl-phosphatidylinositol (GPI) anchor; and
- (ii) a second polypeptide region linked to the first polypeptide region, said second polypeptide region comprising a transmembrane domain from a heterologous protein; or
- a sequence complementary to (a); or (b)
- (c) both (a) and (b)

said molecule being substantially free of nucleic acid molecules not containing (a), (b), or (c).

- The nucleic acid molecule of Claim 1 wherein said chimeric protein has greater than 50% of the complement inhibitory activity of the parent terminal complement inhibitor protein.
- The nucleic acid molecule of Claim 1 wherein the portion of the parent terminal complement inhibitor protein comprises said protein minus amino acid residues downstream of its Ly-6 motif.
- The nucleic acid molecule of Claim 1 wherein the chimeric protein has complement inhibitory activity against human complement.
- A nucleic acid vector comprising the nucleic acid molecule of Claim 1 operatively linked to a second nucleic acid molecule so that a host containing the vector expresses the chimeric protein.
- A recombinant host containing the vector of Claim 4.

- 7. A process for protecting a non-human organ from human complement attack comprising introducing the nucleic acid molecule of Claim 4 into a pluripotent cell capable of producing a non-human transgenic animal and producing the non-human transgenic animal from said cell, whereby the resistance of an organ of said non-human transgenic animal to human complement attack is enhanced.
- 8. Cells isolated from the transgenic animal of Claim 7.
 - 9. A chimeric protein comprising:
- (i) a first polypeptide region comprising a portion of a parent terminal complement inhibitor protein, said portion including a complete Ly-6 motif and not including an operative signal sequence directing the attachment of a glycosyl-phosphatidylinositol (GPI) anchor; and
- (ii) a second polypeptide region linked to the first polypeptide region, said second polypeptide region comprising a transmembrane domain from a heterologous protein.
- 10. The chimeric protein of Claim 9 wherein said protein has greater than 50% of the complement inhibitory activity of the parent terminal complement inhibitor protein.
- 11. The chimeric protein of Claim 9 wherein the portion of the parent terminal complement inhibitor protein comprises said parent protein minus amino acid residues downstream of its Ly-6 motif.
- 12. The chimeric protein of Claim 9 wherein the protein has complement inhibitory activity against human complement.
- 13. A membrane bound terminal complement inhibitor protein which is substantially resistant to the action of lipase enzymes which can cleave a glycosylphosphatidylinositol (GPI) anchor.
- 14. The membrane bound terminal complement inhibitor protein of Claim 13 wherein the protein comprises a Ly-6 motif.

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MGI QGG SVL FGL LLV LAV FCH SGH S LQ CYN CPN PTA DCK MGI QGG SVL FGL LLA LAV FCH SGH S LQ CYN CPN PTT NCK MGI QGG SVL FGL LLV LAV FCH SGH S LQ CYS CPY PTT DCK MGI QGG SVL FGL LLV LAV FCH SGH S LQ CYS CPY PTT QCT MGI QGG SVL FGL LLI LAV FCH SGH S LQ CYS CPY STA RCT MGI QGG SVL FGL LLV LAV FCH SGH S LQ CYS CPY STA RCT MGI QGG SVL FGL LLV LAV FCH SGH S LQ CYS CPL PTM ESM ECT M YIL FTL VLT F-V FCK PIH S LQ CYN CSH STM QCT	TAV NCS SDF DAC LIT KAG LQV YNK CWK FEH CNF NDV TTR LRE NEL TAI NCS SGF DTC LIA RAG LQV YNQ CWK FAN CNF NDI STL LKE SEL TAI NCS SGF DTC LIA RAG LQV YNQ CWK FAN CNF NDI STL LKE SEL MTT NCT SNL DSC LIA KAG SRV YYR CWK FED CTF SRV SNQ LSE NEL TTT NCT SNL DSC LIA KAG SRV YYR CWK FED CTF RQL SNQ LSE NEL AST NCT SNL DSC LIA KAG SGV YYR CWK FDD CSF KRI SNQ LSE TQL TST SCT SNL DSC LIA KAG SGV YYR CWK FDD CSF KRI SNQ LSE TQL TST SCT SNL DSC LIA KAG SGV YYR CWK FDD CSF KRI SNQ LSE TQL	TYY CCK KDL CNF NEO LEN GGT SLS EKT VLL LVT PFL AAA WSL HP QYF CCK EDL CNEQ LEN GGT SLS EKT VLL LVT PLL AAA WCL HP QYF CCK KDL CNF NEQ LEN GGT SLS EKT VVL LVT LLL AAA WCL HP KYY CCK KNL CNF NEA LKN GGT TLS KKT VLL LVI PFL VAA WSL HP KYH CCR ENL CNF NGI LEN GGT TLS KKT VLL LVT PFL AAA WSL HP KYH CCK KNL CNV KEV LEN GGT TLS KKT ILL LVT PFL AAA WSR HP KYH CCK KNL CNV NKG IEN IKR TIS DKA LLL LLA LFL VTA WNF PL
CD59 AGMCIP BABCIP OWMCIP MARCIP SQMCIP	CD59 TAV AGMCIP TAI BABCIP TAI OWMCIP MTT MARCIP TTT SQMCIP AST HVS-15 TST	59 TYY MCIP QYF BCIP QYF MCIP KYY RCIP KYH MCIP KYH
HONONA	HOZODAU	CD AGI BA MAN HV

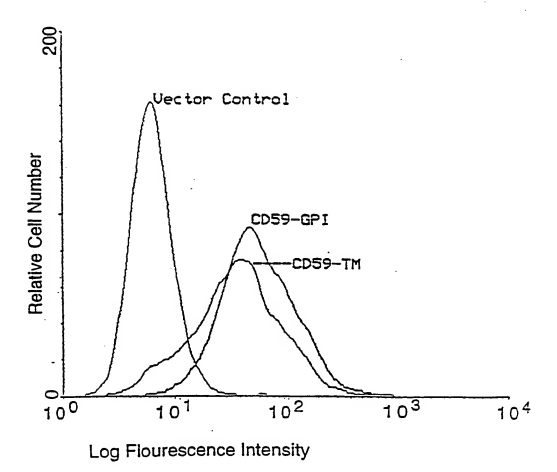
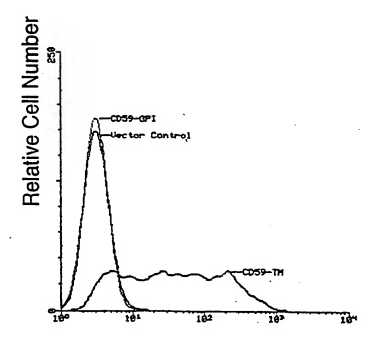
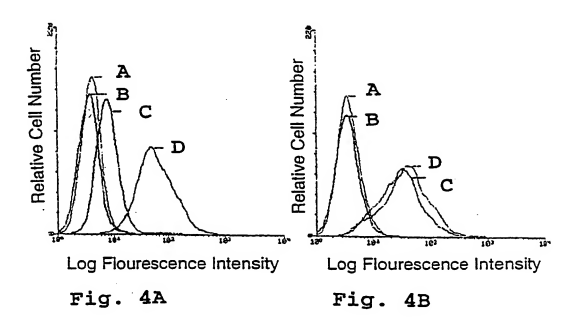


Fig. 2



Log Flourescence Intensity

Fig. 3



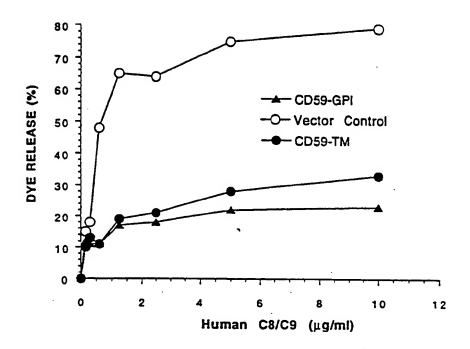


Fig. 5

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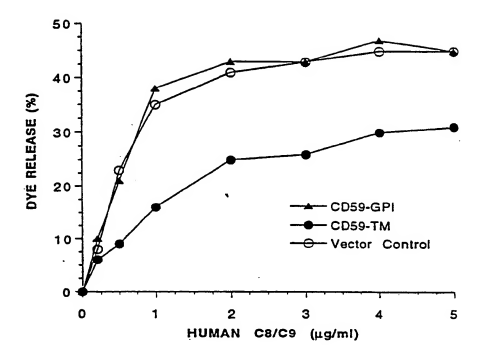


Fig. 6

INTERNATIONAL SEARCH REPORT

International application No. PCT/US95/02944

A. CLASSIFICATION OF SUBJECT MATTER IPC(6) :Please See Extra Sheet.					
US CL :424/93.21; 435/41, 69.1, 69.7, 172.1, 172.3, 240.1, 240.21, 243; 514/44; 530/380; 536/23.4 According to International Patent Classification (IPC) or to both national classification and IPC					
B. FIELDS SEARCHED					
Minimum documentation searched (classification system follow	ed by classification symbols)				
U.S.: 424/93.21; 435/41, 69.1, 69.7, 172.1, 172.3, 240.	1, 240.21, 243; 514/44; 530/380; 536/23.4				
Documentation searched other than minimum documentation to to	Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched				
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)				
Please See Extra Sheet.					
C. DOCUMENTS CONSIDERED TO BE RELEVANT					
Category* Citation of document, with indication, where a	appropriate, of the relevant passages Relevant to claim No.				
HUMAN GENE THERAPY, Volume 2, issued 1991, F.D. 6-8 Ledley, "Clinical considerations in the design of protocols for somatic gene therapy", pages 77-83, see entire document.					
Y THE LANCET, Volume 339, issue Gutierrez et al., "Gene therapy for see entire article.	THE LANCET, Volume 339, issued 21 March 1992, A.A. 6-8 Gutierrez et al., "Gene therapy for Cancer", pages 715-721, see entire article.				
X,P BLOOD, Volume 84, Number 8, issued 15 October 1994, R.P. Rother et al., "Expression of recombinant transmembrane CD59 in paroxysmal nocturnal hemoglobinuria B cells confers resistance to human complement", pages 2604-2611, see entire document.					
X Further documents are listed in the continuation of Box	C. See patent family annex.				
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Date of the actual completion of the international search 30 MAY 1995 Date of mailing of the international search report 07 JUN 1995					
Name and mailing address of the ISA/US Commissioner of Palents and Trademarks Box PCT Washington, D.C. 20231 Authorized officer BRIAN R. STANTON Authorized officer BRIAN R. STANTON					
Facsimile No. (703) 305-3230 Telephone No. (703) 308-0196					

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INTERNATIONAL SEARCH REPORT

International application No.
PCT/US95/02944

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.		
Y	EXPERIMENTAL AND CLINICAL IMMUNOGENETICS, Volume 9, issued 1992, G.T. Venneker et al., "CD59: A molecule involved in antigen presentation as well as downregulation of membrane attack complex", pages 33-47, see entire document.	1-14		
Y	EUROPEAN JOURNAL OF IMMUNOLOGY, Volume 21, issued 1991, L.A. Walsh et al., "Transfection of human CD59 complementary DNA into rat cells confers resistance to human complement", pages 847-850, see entire document.	1-14		
Y	CELL, Volume 73, issued 21 May 1993, Takeda et al., "Deficiency of the GPI anchor caused by a somatic mutation of the PIG-A gene in paroxysmal nocturnal hemoglobinuria", pages 703-711, see entire document.	1-14		
Y	JOURNAL OF EXPERIMENTAL MEDICINE, Volume 174, issued July 1991, D.M. Lublin et al., "Phospholipid-anchored and transmembrane versions of either decay-accelerating factor or membrane cofactor protein show equal efficiency in protection from complement-mediated cell damage", pages 35-44, see entire document.			
Y	DNA AND CELL BIOLOGY, Volume 9, Number 3, issued 1990, R. Sawada et al., "Isolation and expression of the full-length cDNA encoding CD59 antigen of human lymphocytes", pages 213-220, see entire document.	1-14		
Y	SEMINARS IN HEMATOLOGY, Volume 26, Number 3, issued July 1989, B. Rotoli et al., "Paroxysmal nocturnal hemoglobinuria", pages 201-207, see entire document.	1-14		
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INTERNATIONAL SEARCH REPORT

International application No. PCT/US95/02944

			PCT/US95/02944	
A. CLASSIFICATION O IPC (6):	F SUBJECT MATTER:	•		
AO1N 63/00; A61K 35/1 15/09; 15/06, 15/11; C12	4, 38/00; C07H 17/00; C07K 14/00 P 1/00; 21/06	C12N 1/00, 5/00; 5/0	06, 5/22, 7/01, 15/00, 15/0	93,
B. FIELDS SEARCHED Electronic data bases cons	sulted (Name of data base and where	practicable terms used	i):	
Databases: APS, Medline, CA, CAPreviews, Embase, Derwent, Biosis Search Terms:chimer?; protein?; terminal; complement; inhibitor?; ly-6; ly6; glycosyl?; phosphatidylinositol; gpi; transmembran?; anchor; motif; cysteine?; vector?; lysis; mirl; cd59; daf; transgen?; lipase?; rother?/au; rollins?/au; squinto?/au				
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